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Bogle et al.

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(54) TARTRATE SALTS OF 5,8, 14-TRIAZATERACYCLO[10.3.1.02,11 04.9]-HEXADECA-2(11),3,5,7,9-PENTAENE AND PHARMACEUTICAL COMPOSITIONS THEREOF

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- (60) Provisional application No. 60/290,861, filed on May 14, 2001.
- (51) **Int. Cl.**⁷ **C07D 241/36**; A61K 31/50; A61K 31/495
- (52) **U.S. Cl.** **514/252.1**; 514/255.04; 544/343

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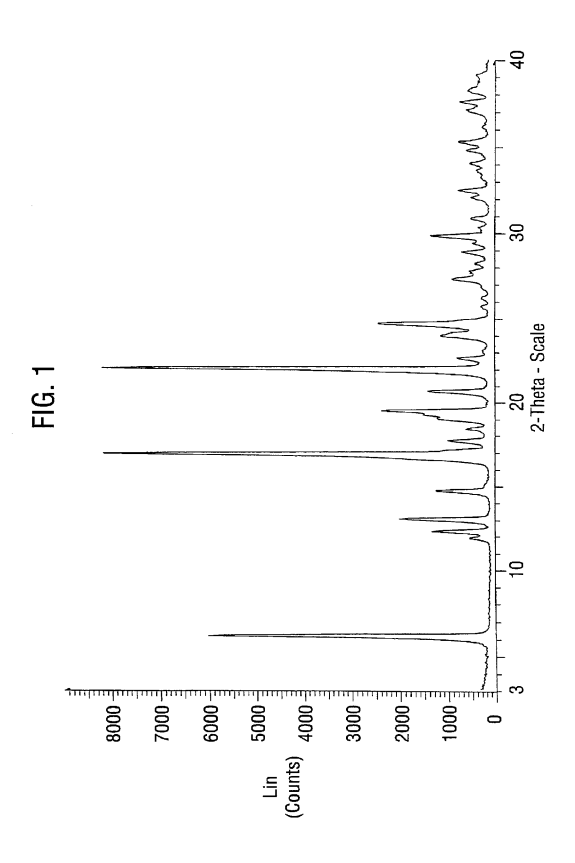
Primary Examiner—Bruck Kifle (74) Attorney, Agent, or Firm—Peter C. Richardson; Lorraine B. Ling; A. David Joran

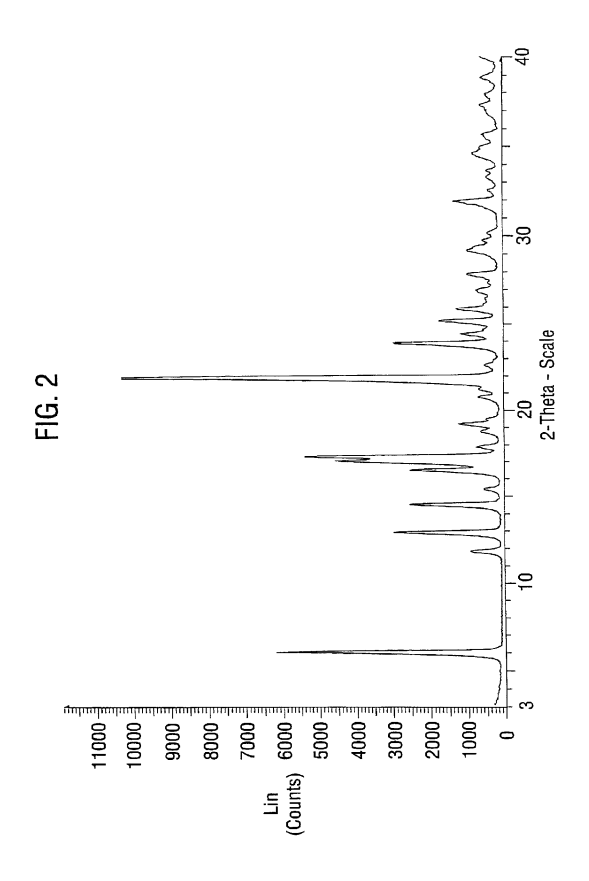
(57) ABSTRACT

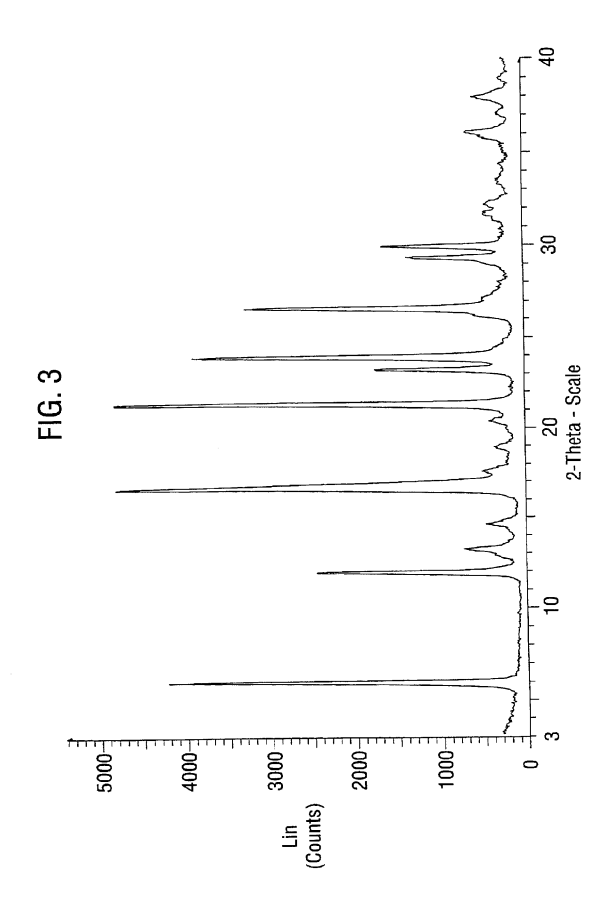
The present invention is directed to the tartrate salts of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene:

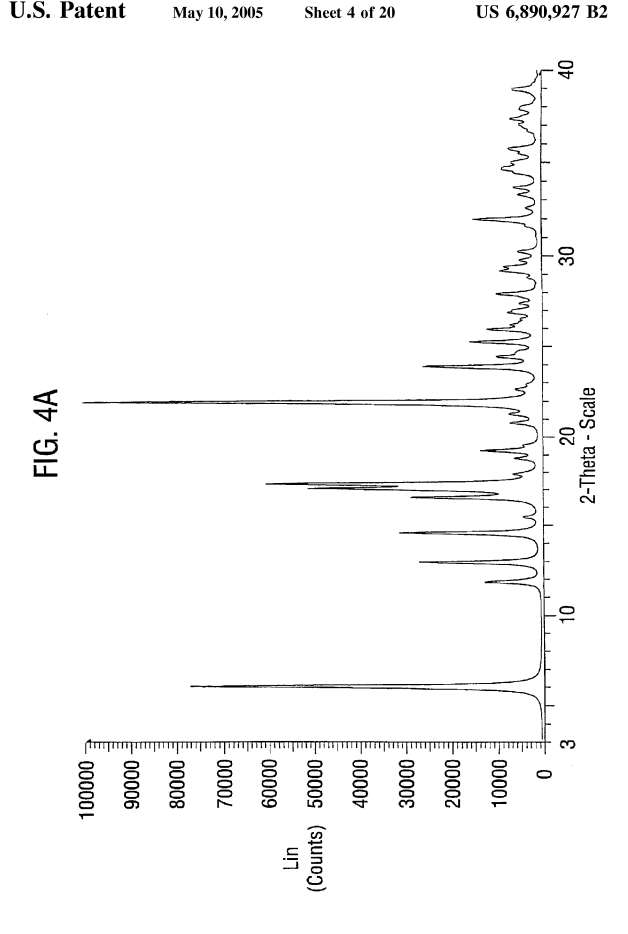
and pharmaceutical compositions thereof. The present invention in particular is directed to the L-tartrate salt, and further to the various polymorphs of the L-tartrate salt, including two distinct anhydrous polymorphs (referred to herein as Forms A and B) and a hydrate polymorph (referred to herein as Form C). In addition, the present invention is also directed to the D-tartrate salt of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene and the various polymorphs thereof; as well as the D,L-tartrate salt thereof and its polymorphs, and the meso-tartrate salt thereof and its polymorphs.

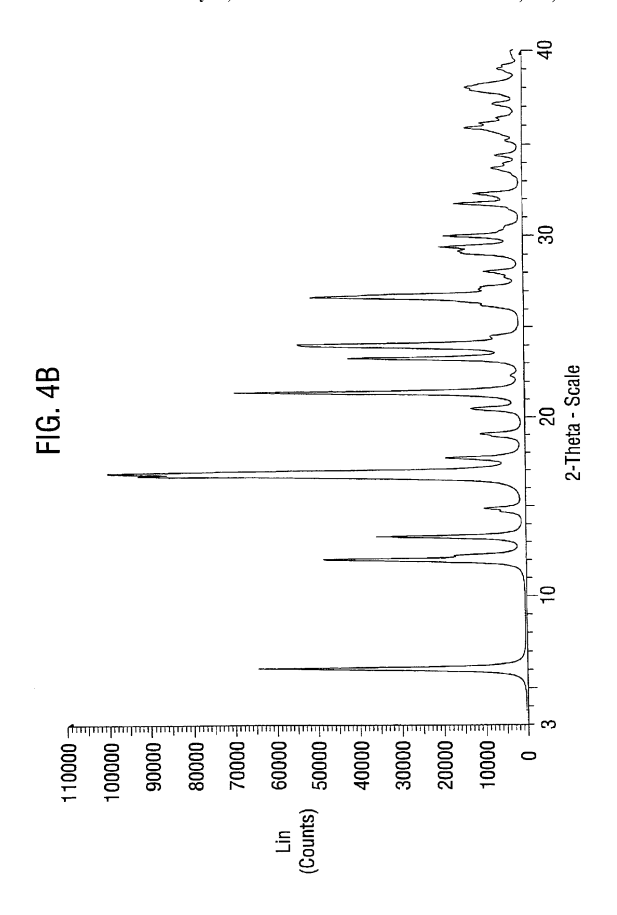
40 Claims, 20 Drawing Sheets

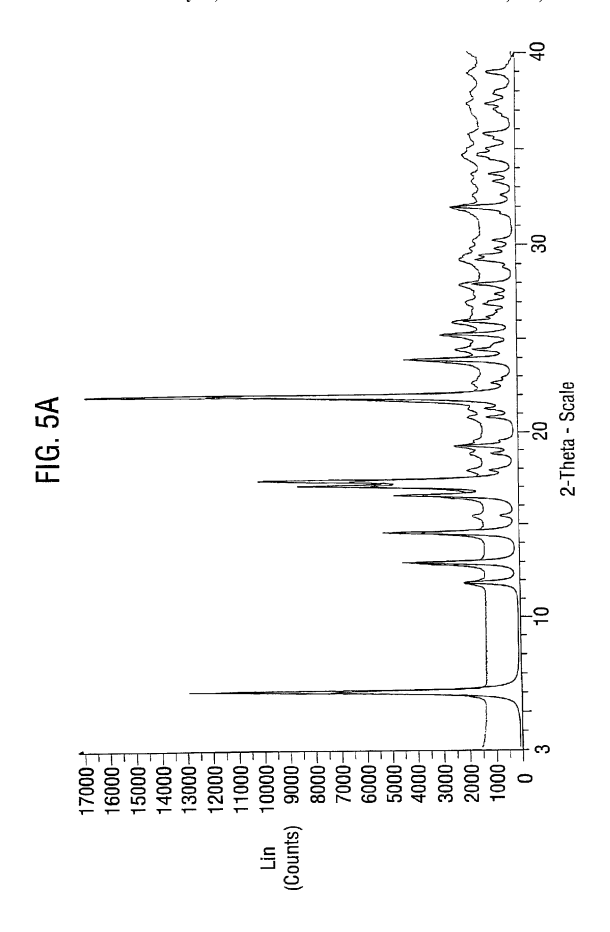


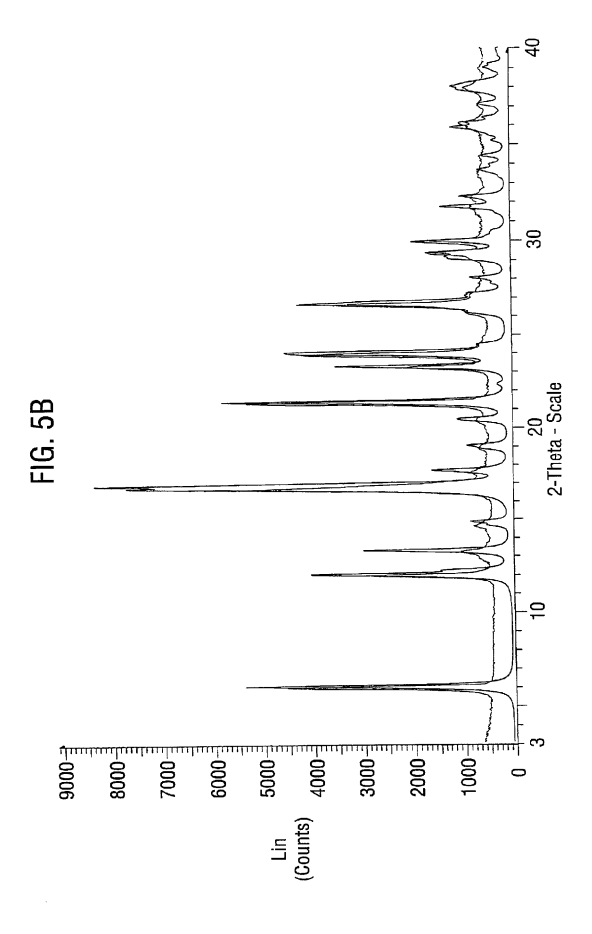


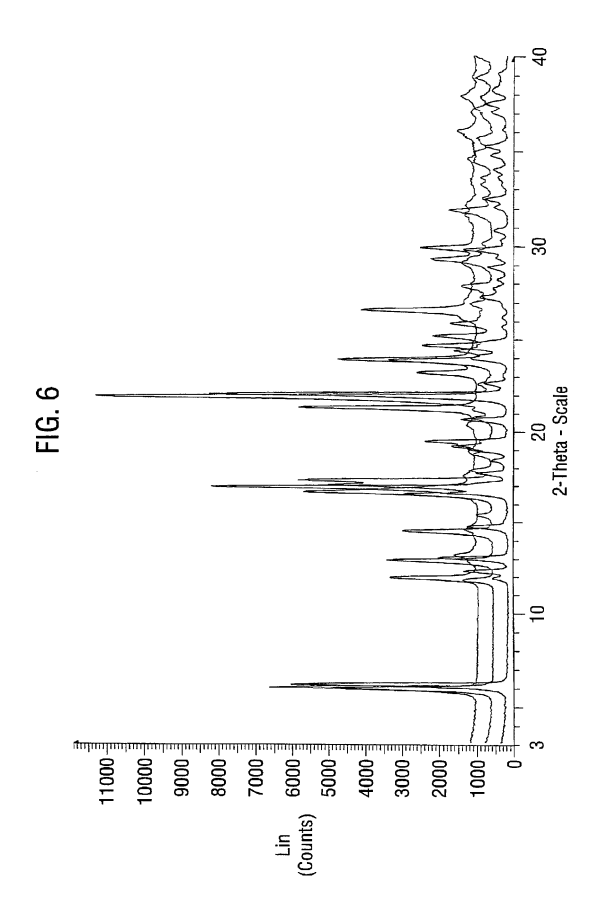


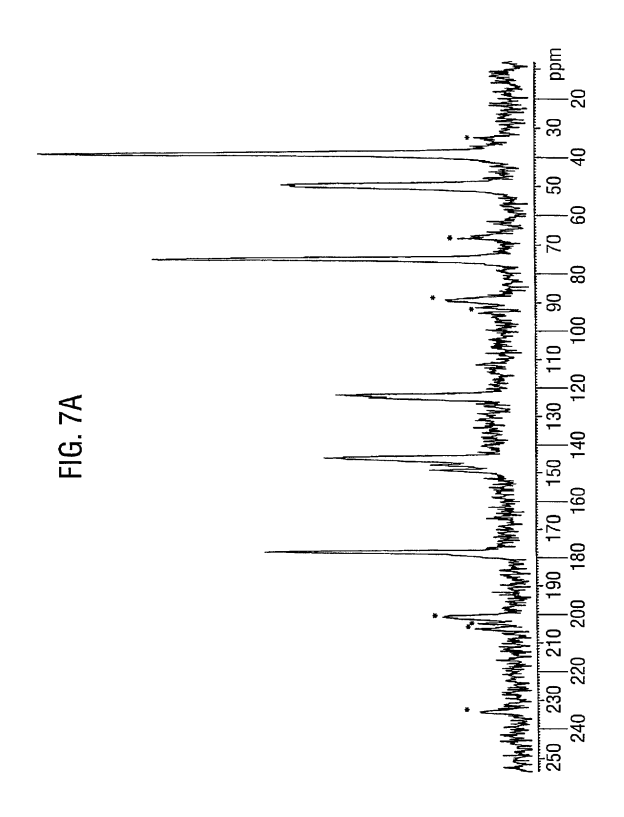


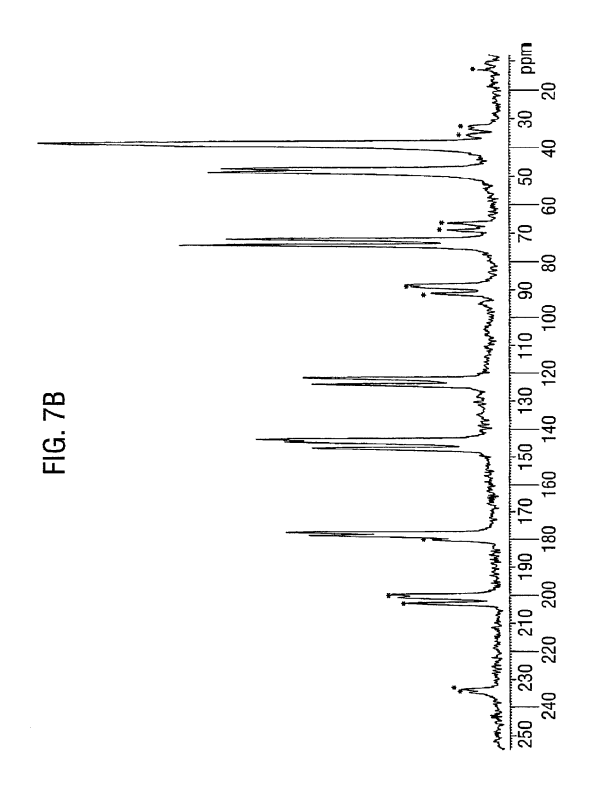












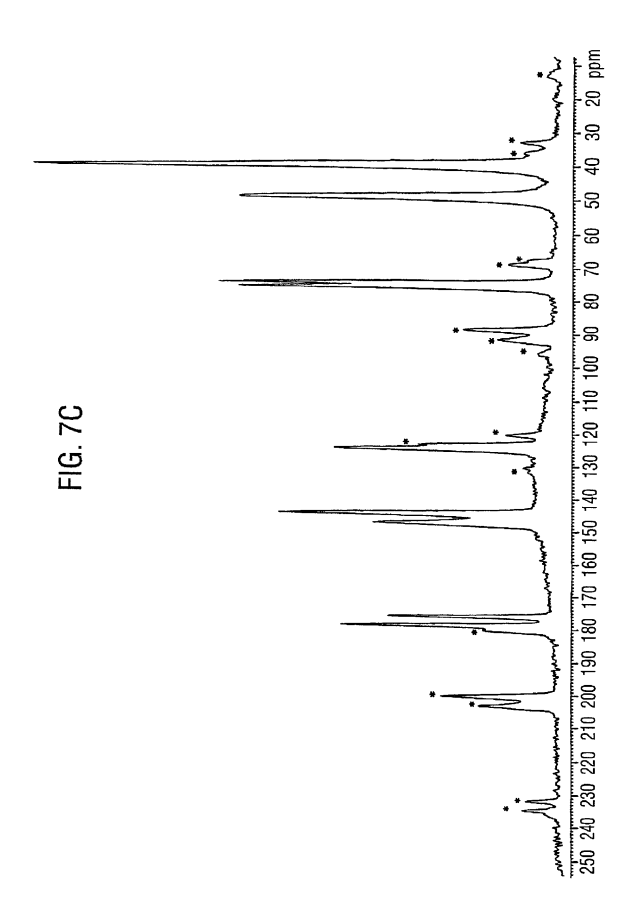


FIG. 8A

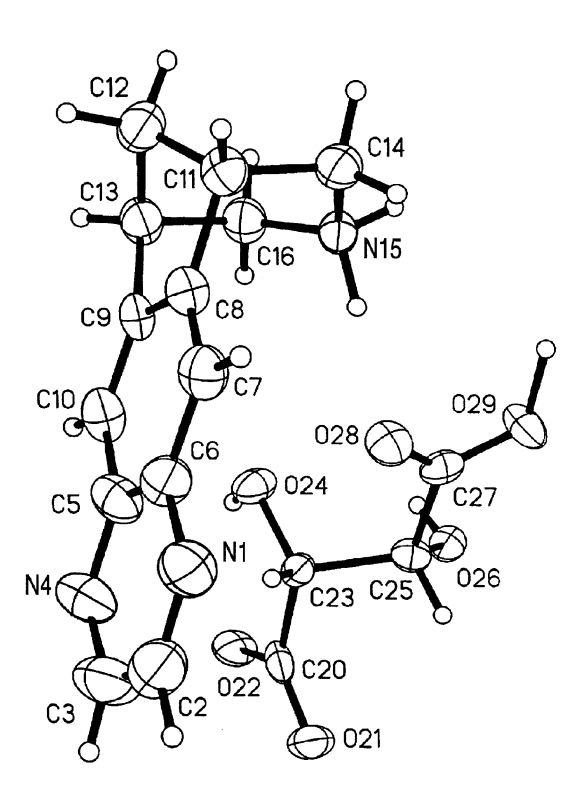
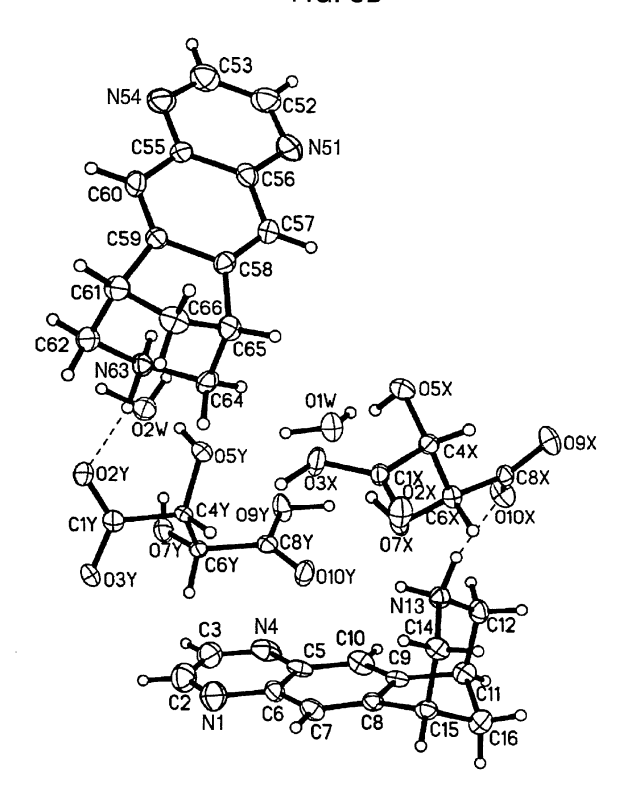
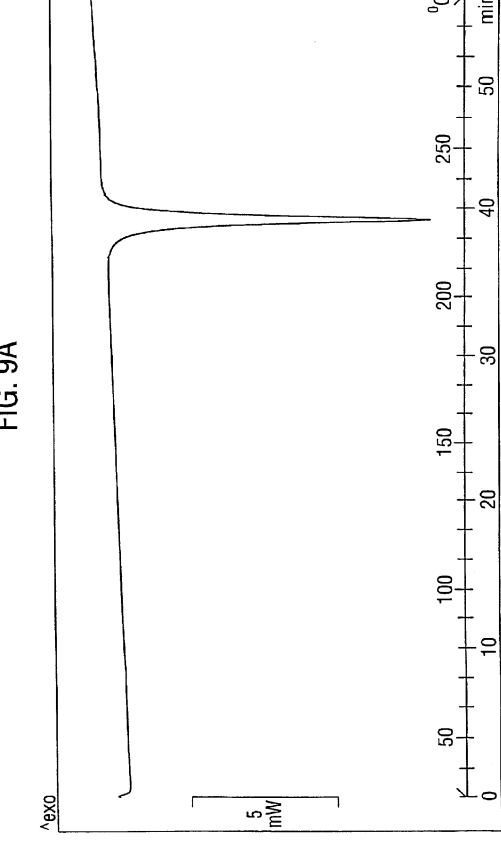
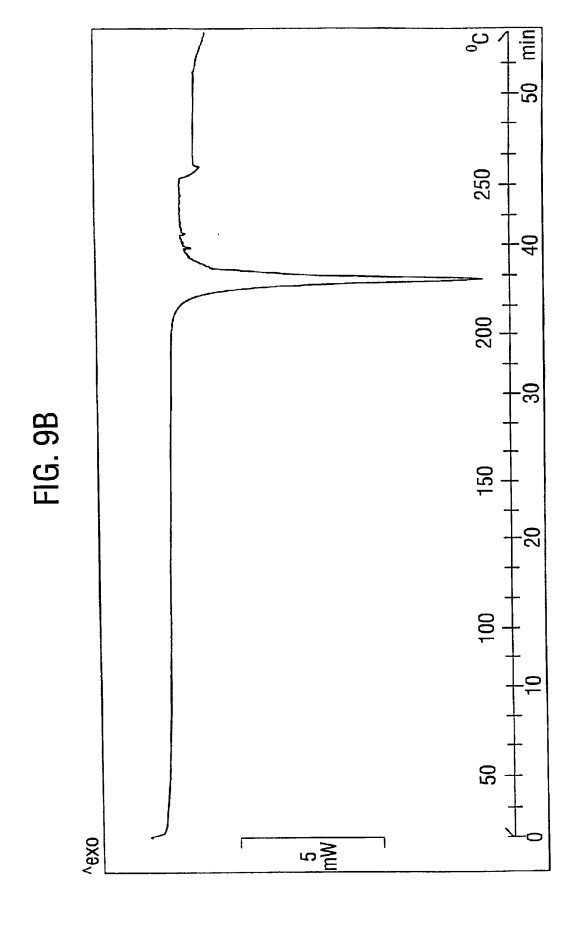


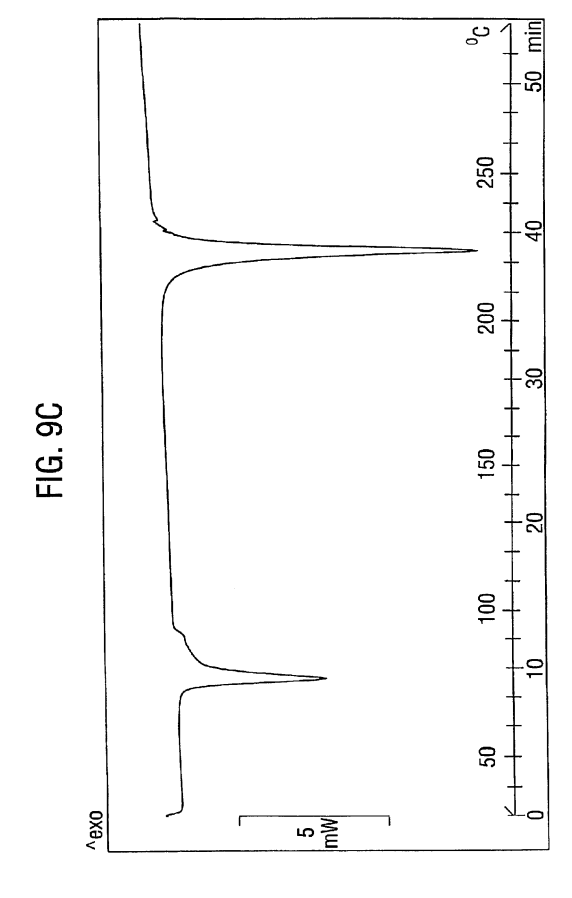
FIG. 8B

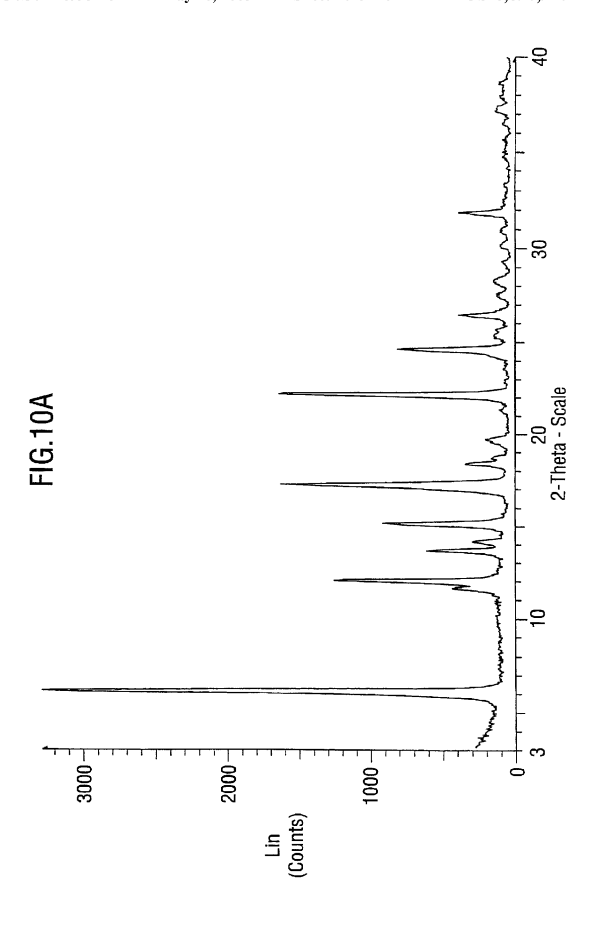


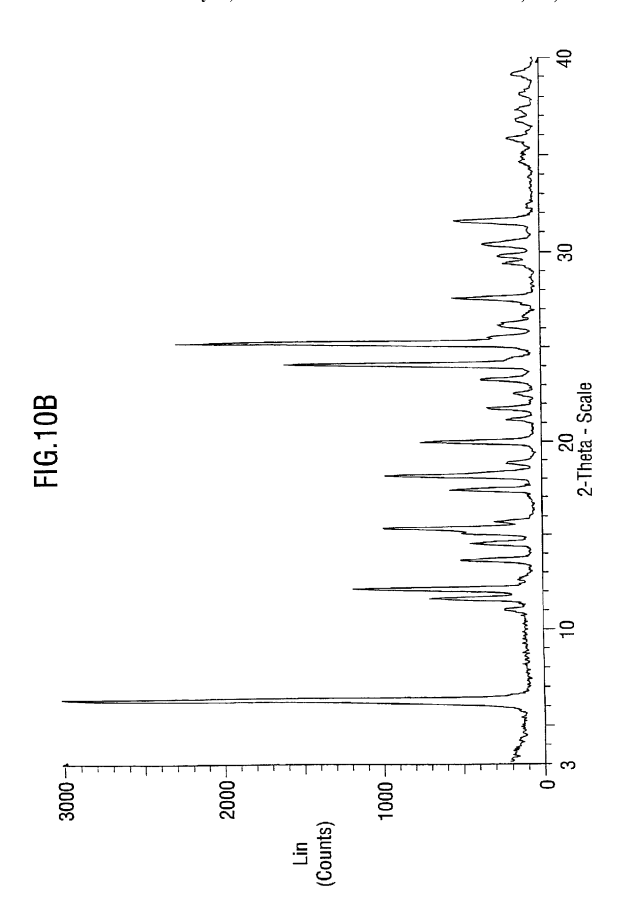


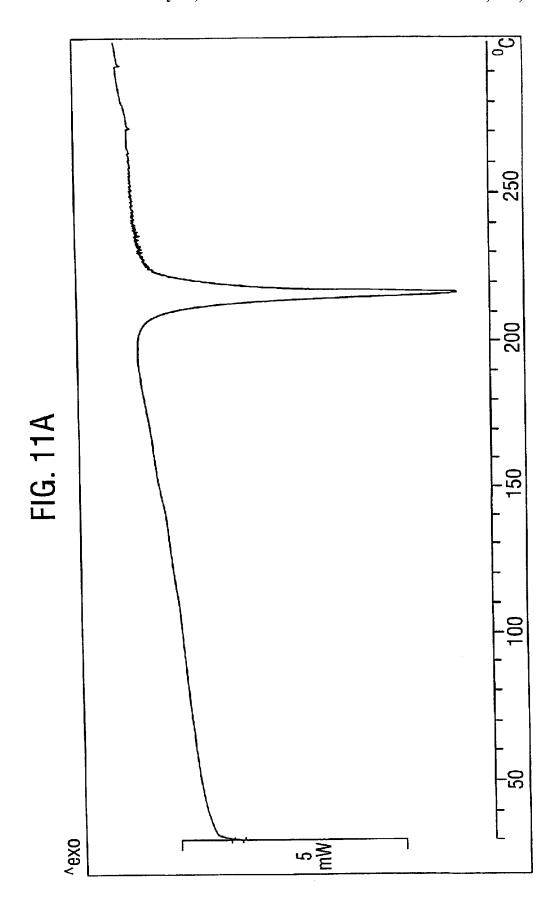


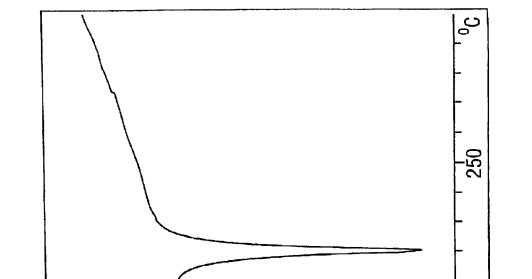
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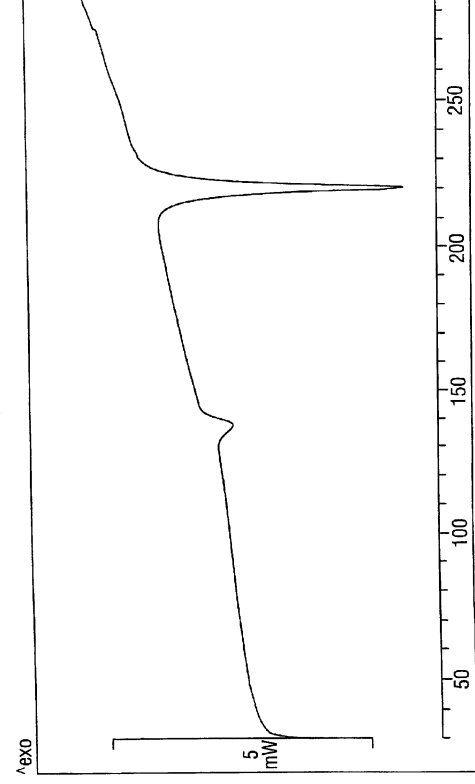












TARTRATE SALTS OF 5.8, 14-TRIAZATERACYCLO[10.3.1.02,11 04.9]-HEXADECA-2(11),3,5,7,9-PENTAENE AND PHARMACEUTICAL COMPOSITIONS THEREOF

This application claims the benefit of U.S. Provisional Application Ser. No. 60/290,861, filed May 14, 2001.

The present invention is directed to the tartrate salts of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, ₁₀ 5,7,9-pentaene:

and pharmaceutical compositions thereof. The present invention in particular is directed to the L-tartrate salt, and further to the various polymorphs of the L-tartrate salt, 20 including two distinct anhydrous polymorphs (referred to herein as Forms A and B) and a hydrate polymorph (referred to herein as Form C). In addition, the present invention is also directed to the D-tartrate salt of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene and the 25 various polymorphs thereof; as well as the D,L-tartrate salt thereof and its polymorphs, and the meso-tartrate salt thereof and its polymorphs.

The compound, 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]hexadeca-2(11),3,5,7,9-pentaene, binds to neuronal nico- 30 drug product formulation use. tinic acetylcholine specific receptor sites and is useful in modulating cholinergic function. This compound is useful in the treatment of inflammatory bowel disease (including but not limited to ulcerative colitis, pyoderma gangrenosum and Crohn's disease), irritable bowel syndrome, spastic 35 dystonia, chronic pain, acute pain, celiac sprue, pouchitis, vasoconstriction, anxiety, panic disorder, depression, bipolar disorder, autism, sleep disorders, jet lag, amyotrophic lateral sclerosis (ALS), cognitive dysfunction, drug/toxin-induced cognitive impairment (e.g., from alcohol, barbiturates, vita- 40 min deficiencies, recreational drugs, lead, arsenic, mercury), disease-induced cognitive impairment (e.g., arising from Alzheimer's disease (senile dementia), vascular dementia, Parkinson's disease, multiple sclerosis, AIDS, encephalitis, trauma, renal and hepatic encephalopathy, hypothyroidism, 45 Pick's disease, Korsakoff's syndrome and frontal and subcortical dementia), hypertension, bulimia, anorexia, obesity, cardiac arrhythmias, gastric acid hypersecretion, ulcers, pheochromocytoma, progressive supramuscular palsy, chemical dependencies and addictions (e.g., dependencies 50 on, or addictions to nicotine (and/or tobacco products), alcohol, benzodiazepines, barbiturates, opioids or cocaine), headache, migraine, stroke, traumatic brain injury (TBI), obsessive-compulsive disorder (OCD), psychosis, Huntingschizophrenia, multi-infarct dementia, age-related cognitive decline, epilepsy, including petit mal absence epilepsy, attention deficit hyperactivity disorder (ADHD), Tourette's Syndrome, particularly, nicotine dependency, addiction and withdrawal; including use in smoking cessation therapy.

The tartrate salts of this invention may also be used in a pharmaceutical composition in combination with an antidepressant such as, for example, a tricyclic antidepressant or a serotonin reuptake inhibiting antidepressant (SRI), in order to treat both the cognitive decline and depression associated 65 with AD, PD, stroke, Huntington's chorea or traumatic brain injury (TBI); in combination with muscarinic agonists in

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order to stimulate both central muscarinic and nicotinic receptors for the treatment, for example, of ALS, cognitive dysfunction, age-related cognitive decline, AD, PD, stroke, Huntington's chorea and TBI; in combination with neurotrophic factors such as NGF in order to maximize cholinergic enhancement for the treatment, for example, of ALS, cognitive dysfunction, age-related cognitive decline, AD, PD stroke, Huntington's chorea and TBI; or in combination with agents that slow or arrest AD such as cognition enhancers, amyloid aggregation inhibitors, secretase inhibitors, tau kinase inhibitors, neuronal anti-inflammatory agents and estrogen-like therapy.

Compounds that bind to neuronal nicotinic receptor sites, including 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ 15 hexadeca-2(11),3,5,7,9-pentaene, and its hydrochloride salt, are referred to in WO 99/35131, published Jul. 15, 1999 (corresponding to U.S. Ser. No. 09/402,010, filed Sep. 28, 1999 and Ser. No. 09/514,002, filed Feb. 25, 2000). The foregoing applications, owned in common with the present application and incorporated herein by reference in their entirety, generically recite pharmaceutically acceptable acid addition salts for the compounds referred to therein.

The L-tartrate salt of the present invention exhibits properties, including those of high solid-state stability and compatibility with certain drug product formulation excipients, that render it superior to previously known salts of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11), 3,5,7,9-pentaene. Further, the D-tartrate and D,L-tartrate salts exhibit properties that also make them appropriate for

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a powder X-ray diffraction of the anhydrous L-tartrate salt Form A of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIG. 2 is the powder X-ray diffraction of the anhydrous L-tartrate salt Form B of 5,8,14-triazatetra-cyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIG. 3 is the powder X-ray diffraction of the L-tartrate salt hydrate Form C of 5,8,14-triazatetra-cyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIG. 4A is the calculated powder X-ray diffraction pattern of the anhydrous Form B L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta). FIG. 4B is the calculated powder X-ray diffraction pattern of the Form C L-tartrate salt hydrate of 5,8,14-triazatetra-cyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIG. 5A is the calculated powder X-ray diffraction pattern ton's chorea, tardive dyskinesia, hyperkinesia, dyslexia, 55 (lower trace) laid over the observed X-ray diffraction pattern (upper trace) for the anhydrous Form B L-tartrate salt of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta). FIG. 5B is the calculated powder X-ray diffraction pattern (lower trace) laid over the observed X-ray diffraction pattern (upper trace) for the Form C L-tartrate salt hydrate of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

> FIG. 6 is the overlay of the powder X-ray diffraction patterns of the Form A (lower trace), Form B (middle trace) and Form C (upper trace) L-tartrate salts of 5,8,14-

triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIGS. 7A, 7B and 7C are the solid state ¹³C NMR spectra of the L-tartrate salts of 5,8,14-triazatetra-cyclo ⁵ [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene Forms A, B and C, respectively, as measured by cross-polarization magic angle spinning (CPMAS) at 295 K on a Bruker 7 mm wide-bore magic angle spinning (WB MAS) probe positioned in a Bruker Avance DRX 500 MHz NMR Spectrometer. Peaks marked with asterisks (*) are spinning sidebands which are displaced at multiples of the spinning frequencies along both sides of the real peaks (centerbands).

FIG. **8A** is the X-ray crystal structure (absolute configuration) for the anhydrous Form B L-tartrate salt of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene. FIG. **8B** is the X-ray crystal structure (absolute configuration) for the Form C L-tartrate salt hydrate of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene.

FIG. **9A**, **9B** and **9**C are the differential scanning calorimetric traces for the L-tartrate salts Forms A, B and C, respectively, of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene.

FIG. **10**A and **10**B are the powder X-ray diffraction patterns of the D,L-tartrate salt Forms X and Y, respectively, of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11), 3,5,7,9-pentaene (y axis is linear counts per second; X in degrees 2 theta).

FIGS. 11A and 11B are the differential scanning calorimetric traces for the D,L-tartrate salts Forms X and Y, respectively, of 5,8,14-triazatetra-cyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene.

SUMMARY OF THE INVENTION

The present invention relates to the tartrate salts of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene. The tartrate salts of the invention include the L-tartrate, D-tartrate, D,L-tartrate and meso-tartrate salts.

In particular, the present invention relates to the L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2 (11),3,5,7,9-pentaene.

In one embodiment of the invention, the L-tartrate of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene is the anhydrous L-tartrate salt, referred to herein as Form A. The L-tartrate Form A is characterized by the principal x-ray diffraction pattern peaks expressed in terms of 2θ and d-spacings as measured with copper radiation (within the margins of error indicated):

Angle 2θ (±0.2)	d-value (Å) (±0.2)	<u> </u>
6.1	14.5	— ,
12.2 13.0	7.2 6.8	
14.7	6.0	
16.8	5.3	
19.4 21.9	4.6 4.1	(
24.6	3.6	

The L-tartrate crystal Form A is characterized in that it has a onset of melt at about 223° C. as measured by differential 65 scanning calorimetry at a heating rate of 5 degrees per minute. The L-tartrate Form A is also characterized in that

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when examined by solid state ¹³C NMR cross-polarization magic angle spinning techniques, it exhibits the following principal resonance peaks (±0.1 ppm) downfield from 100 ppm (adamantane standard 29.5 ppm): 178.4, 149.3, 147.4, 145.1, and 122.9 ppm.

In another embodiment of the invention, the L-tartrate of 5,8,14-triazatetracyclo[$10.3.1.0^{2,11}.0^{4,9}$]-hexadeca-2(11),3, 5,7,9-pentaene is another anhydrous L-tartrate salt polymorph, referred to herein as Form B. The L-tartrate salt Form B is characterized by the principal x-ray diffraction pattern peaks expressed in terms of 2θ and d-spacings as measured with copper radiation (within the margins of error indicated):

Angle 2θ (±0.2)	d-value (Å) (±0.2)
5.9	15.0
12.8	6.9
14.4	6.1
15.3	5.8
16.9	5.2
17.2	5.2
21.8	4.1
23.8	3.7
25.1	3.5

The L-tartrate salt Form B has a single crystal x-ray structure (absolute configuration) as set forth in FIG. 8A. Further, the Form B forms orthorhombic crystals belonging to the P2(1)2(1)2(1) space group. Form B is further characterized in having an onset of melting at about 215° C. as measured by differential scanning calorimetry at a heating rate of 5 degrees per minute. Further, Form B of the invention is also characterized in having an aqueous solubility of about 156 mg/ml and a native pH of about 3.3 in aqueous solution. In addition, Form B has a hygroscopicity of approximately 0.2% at 90% relative humidity.

The L-tartrate Form B is also characterized in that when examined by solid state ¹³C NMR cross-polarization magic angle spinning techniques, it exhibits the following principal resonance peaks (±0.1 ppm) downfield from 100 ppm (adamantane standard 29.5 ppm): 179.2, 178.0, 147.4, 145.2, 144.4, 124.8 and 122.5 ppm.

In another embodiment of the invention, the L-tartrate of 5.8.14-triazatetracyclo[$10.3.1.0^{2.11}.0^{4.9}$]-hexadeca-2(11),3, 5.7.9-pentaene is the hydrate L-tartrate salt, referred to herein as Form C. The L-tartrate Form C is characterized by the principal x-ray diffraction pattern peaks expressed in terms of 2θ and d-spacings as measured with copper radiation (within the margins of error indicated):

<u> </u>	Angle 2θ (±0.2)	d-value (Å) (±0.2)	
55 —	5,9	15.1	
	11.8	7.5	
	16.5	5.4	
	21.2	4.2	
	23.1	3.8	
60	23.8	3.7	
00	26.5	3.4	
			_

The hydrate L-tartrate crystal Form C has a single crystal structure as set forth in FIG. 8B. Further, the hydrate Form C forms monoclinic crystals belonging to the P2(1) space group. Form C is further characterized in having an onset of a solid-solid transition at about 72° C. and an onset of

melting transition at about 220° C. Because Form B converts to the hydrate Form C upon contact with 100% relative humidity, Form C has the same aqueous solubility as Form B.

The L-tartrate Form C is also characterized in that when 5 examined by solid state ¹³C NMR cross-polarization magic angle spinning techniques, it exhibits the following principal resonance peaks (±0.1 ppm) downfield from 100 ppm (adamantane standard 29.5 ppm): 179.0, 176.1, 147.5, 144.5 and 124.6 ppm.

A further embodiment of the invention is directed to the D-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene. In particular, the present invention is directed to the three D-tartrate salt polymorphs (referred to here as Forms A', B' and C') which exhibit the ¹⁵ same x-ray diffraction characteristics, hygroscopicity, water content and thermal characteristics as Forms A, B and C of the L-tartrate salt.

In another embodiment, the present invention relates to the D,L-tartrate salt of 5,8,14-triazatetracyclo ²⁰ [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene, and in particular, two polymorphs, an anhydrous form (herein referred to as Form X) and a hydrate form (herein referred to as Form Y).

The D,L-tartrate Form X is characterized by the principal 25 x-ray diffraction pattern peaks expressed in terms of 2θ and d-spacings as measured with copper radiation (within the margins of error indicated):

Angle 2θ (±0.2)	d-value (Å) (±0.2)
6.0	14.6
11.9	7.4
15.0	5.9
17.1	5.2
22.1	4.0
24.5	3.6

The D,L-tartrate Form X is further characterized in having 40 an onset of a melting transition at about 212° C.

The D,L-tartrate Form Y is characterized by the principal x-ray diffraction pattern peaks expressed in terms of 2θ and d-spacings as measured with copper radiation (within the margins of error indicated):

Angle 2θ (±0.2)	d-value (Å) (±0.2)
6.2	14.2
12.0	7.4
15.2	5.8
18.1	4.9
24.0	3.7
25.1	3.5

The D,L-tartrate Form Y is further characterized in having an onset of a solid-solid transition at about 131° C. and an onset of melting transition at about 217° C.

Another embodiment of the invention relates to a pharmaceutical composition comprising at least one of polymorphic Forms A, B or C, preferably Form B, of the L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2 (11),3,5,7,9-pentaene and a pharmaceutically acceptable carrier or excipient, for use in the treatment of inflammatory 65 bowel disease (including but not limited to ulcerative colitis, pyoderma gangrenosum and Crohn's disease), irritable

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bowel syndrome, spastic dystonia, chronic pain, acute pain, celiac sprue, pouchitis, vasoconstriction, anxiety, panic disorder, depression, bipolar disorder, autism, sleep disorders, jet lag, amyotrophic lateral sclerosis (ALS), cognitive dysfunction, drug/toxin-induced cognitive impairment (e.g., from alcohol, barbiturates, vitamin deficiencies, recreational drugs, lead, arsenic, mercury), disease-induced cognitive impairment (e.g., arising from Alzheimer's disease (senile dementia), vascular dementia, Parkinson's disease, multiple sclerosis, AIDS, encephalitis, trauma, renal and hepatic encephalopathy, hypothyroidism, Pick's disease, Korsakoff's syndrome and frontal and subcortical dementia), hypertension, bulimia, anorexia, obesity, cardiac arrhythmias, gastric acid hypersecretion, ulcers, pheochromocytoma, progressive supramuscular palsy, chemical dependencies and addictions (e.g., dependencies on, or addictions to nicotine (and/or tobacco products), alcohol, benzodiazepines, barbiturates, opioids or cocaine), headache, migraine, stroke, traumatic brain injury (TBI), obsessive-compulsive disorder (OCD), psychosis, Huntington's chorea, tardive dyskinesia, hyperkinesia, dyslexia, schizophrenia, multi-infarct dementia, age-related cognitive decline, epilepsy, including petit mal absence epilepsy, attention deficit hyperactivity disorder (ADHD), and Tourette's Syndrome. Another more preferred embodiment of the invention is wherein the pharmaceutical composition is useful in the treatment of nicotine dependency, addiction and withdrawal; most preferably, for use in smoking cessation therapy.

The present invention further relates to pharmaceutical compositions for the uses described in the foregoing paragraph comprising any one of the D-tartrate salt of, the D,L-tartrate salt of, or the meso-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene.

The present invention further relates to a method of treating inflammatory bowel disease (including but not limited to ulcerative colitis, pyoderma gangrenosum and Crohn's disease), irritable bowel syndrome, spastic dystonia, chronic pain, acute pain, celiac sprue, pouchitis, vasoconstriction, anxiety, panic disorder, depression, bipolar disorder, autism, sleep disorders, jet lag, amyotrophic lateral sclerosis (ALS), cognitive dysfunction, drug/toxin-induced cognitive impairment (e.g., from alcohol, barbiturates, vita-45 min deficiencies, recreational drugs, lead, arsenic, mercury), disease-induced cognitive impairment (e.g., arising from Alzheimer's disease (senile dementia), vascular dementia, Parkinson's disease, multiple sclerosis, AIDS, encephalitis, trauma, renal and hepatic encephalopathy, hypothyroidism, 50 Pick's disease, Korsakoff's syndrome and frontal and subcortical dementia), hypertension, bulimia, anorexia, obesity, cardiac arrhythmias, gastric acid hypersecretion, ulcers, pheochromocytoma, progressive supramuscular palsy, chemical dependencies and addictions (e.g., dependencies on, or addictions to nicotine (and/or tobacco products), alcohol, benzodiazepines, barbiturates, opioids or cocaine), headache, migraine, stroke, traumatic brain injury (TBI), obsessive-compulsive disorder (OCD), psychosis, Huntington's chorea, tardive dyskinesia, hyperkinesia, dyslexia, schizophrenia, multi-infarct dementia, age-related cognitive decline, epilepsy, including petit mal absence epilepsy, attention deficit hyperactivity disorder (ADHD), and Tourette's Syndrome comprises administering to a subject in need of treatment a therapeutically effective amount of any of Forms A, B or C of the L-tartrate salt of 5,8,14triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9pentaene, preferably Form B. Another more preferred

embodiment of the invention relates to a method of treatment for nicotine dependency, addiction and withdrawal, in particular for use in smoking cessation therapy activity, comprising the administration of any of Forms A, B or C of the L-tartrate salt of 5,8,14-triazatetracyclo 5 [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene, preferably Form B, to a subject in need thereof.

The present invention further relates to methods of treatment described in the foregoing paragraph comprising the administration of any of the D-tartrate salt, the D,L-tartrate salt or the meso-tartrate salt of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene to a subject in need thereof.

The term "treating" as used herein, refers to, and includes, reversing, alleviating, inhibiting the progress of, or preventing a disease, disorder or condition, or one or more symptoms thereof; and the term "treatment" refers to the act of treating, as defined above.

The invention also relates to a process for the preparation of the Form A of L-tartrate salt of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene comprising the steps of

- (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with between 1 and 2 equivalents of L-tartaric acid; and
- (ii) collecting the crystals formed.

A preferred embodiment of this invention relates to the above process wherein 1.1 equivalents of L-tartaric acid is employed and the tartaric acid is added to a solution containing the free base. A preferred mode of practicing this process is wherein the contact step is allowed to proceed for less than 2 hours. A more preferred embodiment of this invention relates to the above process wherein the contact step (i.e., step "(i)" above) is allowed to proceed above 45° C. Another preferred embodiment of this invention relates to the above process wherein the suitable solvent is selected from the group consisting of a (C_1-C_6) alkyl alcohol, a (C_1-C_6) alkyl ketone or a (C_1-C_6) alkyl ether, acetonitrile and (C_1-C_6) alkyl esters (e.g., ethyl acetate, isopropyl acetate, etc.). More preferably, the suitable solvent is ethanol or methanol.

The invention further relates to a process for the preparation of Form A' of the D-tartrate salt comprising steps (i) and (ii) referred to above for making Form A of the L-tartrate salt, but using D-tartaric acid in step (i) in place of L-tartaric acid.

The invention also relates to a process for the preparation of Form B of L-tartrate salt of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene comprising the steps of:

- (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with about 1 to about 2.3 equivalents of L-tartaric acid; and
- (ii) collecting the crystals formed.

A preferred embodiment of this invention relates to the above process wherein about 1.1 to about 2.2 equivalents, more preferably 1.1 equivalents, of L-tartaric acid is employed and the free base in solution is added to a solution 60 containing L-tartaric acid. A preferred mode of practicing this process is wherein the contact step is allowed to proceed for a minimum of 1 hours; more preferably, for at least 2 hours; most preferably, longer than 12 hours. A preferred embodiment is wherein the suitable solvent is selected from 65 the group consisting of a (C_1-C_6) alkyl alcohol, a (C_1-C_6) alkyl ketone or a (C_1-C_6) alkyl ether, acetonitrile and

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 (C_1-C_6) alkyl esters (e.g., ethyl acetate, isopropyl acetate, etc.). More preferably, the suitable solvent is methanol or ethanol, most preferably methanol.

The invention further relates to a process for the preparation of Form B' of the D-tartrate salt comprising steps (i) and (ii) referred to above for making Form B of the L-tartrate salt, but using D-tartaric acid in step (i) in place of L-tartaric acid.

Another aspect of the present invention relates to a process for the preparation of the Form C of the L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2 (11),3,5,7,9-pentaene comprising the steps of:

- (i) contacting either of Form A or Form B of the L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene with water; and
- (ii) collecting the crystals formed.

A preferred embodiment of this invention relates to the above process wherein the contacting of step (i) comprises slurrying either of Forms A or B with water with subsequent addition of an organic solvent to promote precipitation of the Form C product. A more preferred embodiment of the process is wherein the organic solvent use to promote precipitation is methanol, ethanol or acetonitrile.

The invention further relates to a process for the preparation of Form C' of the D-tartrate salt comprising steps (i) and (ii) referred to above for making Form C of the L-tartrate salt but using Forms A' or B' of the D-tartrate salt in step (i) in place of Forms A or B of the L-tartrate salt.

The present invention further relates to a process for the preparation of Form X of the D,L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene comprising the steps of:

- (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with about 1 to about 2.3 equivalents of D,L-tartaric acid; and
- (ii) collecting the crystals formed.

A preferred embodiment of this invention relates to the above process wherein about 2.2 equivalents of D,L-tartaric acid is employed and the free base in solution is added to a solution containing D,L-tartaric acid. A preferred mode of practicing this process involves allowing the contact step to proceed for a minimum of 2 hours; more preferably, for at least 12 hours; and most preferably, at least 24 hours.

Another preferred embodiment of this invention relates to the above process for preparing Form X wherein the suitable solvent is anhydrous or nearly anhydrous and is selected from the group consisting of a (C_1-C_6) alkyl alcohol, a (C_1-C_6) alkyl ketone or a (C_1-C_6) alkyl ether, acetonitrile and (C_1-C_6) alkyl esters (e.g., ethyl acetate, isopropyl acetate, etc.). More preferably, the suitable solvent is ethanol.

The present invention further relates to a process for the preparation of Form Y of the D,L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene comprising the steps of:

- (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with about 1 to about 2.3 equivalents of D,L-tartaric acid; and
- (ii) collecting the crystals formed.

A preferred embodiment of this invention relates to the above process wherein about 2.2 equivalents of D,L-tartaric acid is employed and the free base in solution is added to a solution containing D,L-tartaric acid. A preferred mode of practicing this process involves allowing the contact step to

proceed for a minimum of 2 hours; more preferably, for at least 12 hours; most preferably, for at least 24 hours.

Another preferred embodiment of this invention relates to the above process for preparing Form Y wherein the suitable solvent is selected from the group consisting of a (C_1-C_6) 5 alkyl alcohol, a (C_1-C_6) alkyl ketone or a (C_1-C_6) alkyl ether, acetonitrile and (C_1-C_6) alkyl esters (e.g., ethyl acetate, isopropyl acetate, etc.) admixed with water. More preferably, the suitable solvent is ethanol admixed with water; most preferably, 20% aqueous ethanol.

DETAILED DESCRIPTION OF THE INVENTION

The compound, 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene is a nicotinic partial agonist for the treatment of a number of CNS diseases, disorders and conditions including, in particular, nicotine dependency, addiction and withdrawal.

Although in general the salts of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene are all crystalline, the majority of such salts are so significantly hygroscopic as to render them poor candidates for pharmaceutical formulation use. The L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene is very slightly hygroscopic, has high aqueous solubility and is high melting. These characteristics, combined with its relative inertness towards common excipients, make it highly suitable for pharmaceutical formulation use. The D-tartrate salt, the D,L-tartrate salt and the meso-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2 (11),3,5,7,9-pentaene also exhibit favorable characteristics.

The L-tartrate salt exists as three possible forms: two anhydrous forms and one hydrate form. Of the two anhydrous forms, Form A and Form B, Form A is the kinetic 35 polymorph, which will convert under appropriate conditions to the thermodynamically favored Form B. The hydrate L-tartrate salt Form C is a monohydrate and is relatively stable under ambient conditions. It will maintain its one equivalent of water under vacuum at moderate temperatures 40 for at least a day (e.g., for 24 hours in a 45° C. vacuum oven), but eventually over time (i.e., 48 hours or more) will lose water and convert to the anhydrous Form B. Form B is the most stable of the polymorphs at low humidity. Accordingly, Form B would appear to be the most appro- 45 priate and most stable polymorph of the L-tartrate salts of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3, 5,7,9-pentaene for pharmaceutical formulation use.

As noted above, Form A is the anhydrous kinetic polymorph, which converts under appropriate conditions to 50 the thermodynamically-favored Form B. Form A is obtainable from a synthesis involving, e.g., contacting the free base of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2 (11),3,5,7,9-pentaene with approximately one equivalent of L-tartaric acid in methanol or ethanol, allowing little or no 55 time for equilibration. Form A is observed as the resulting product initially from the combination of the 5,8,14triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9pentaene free base and L-tartaric acid, but Form B begins to form on continued or prolonged agitation of the reaction 60 mixture. The rate of formation of Form B may be accelerated by using at least a two-fold or more stoichiometric excess of L-tartaric acid (i.e., faster with 2.2 equivalents of L-tartaric acid present than with only 1.1 equivalents) and allowing the reaction to proceed for longer than two hours, preferably for 65 at least a day or more. Conversion to Form B is ordinarily complete after about 5 hours using 2.2 equivalents. In

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contrast, the conversion may require more than 20 hours using 1.1 equivalents. In any case, conversion to Form B is usually complete under most conditions after 48 hours at 20–25° C.

The temperature of the L-tartrate salt formation reaction also influences whether Form A or Form B is isolated, since Forms A and B appear to be thermally interconvertable. Running the salt formation reaction above 45° C. give Form A. Conversely, formation of the salt below 45° C. results in the formation of predominantly Form B. Also, stirring Form A in methanol below 40° C. results in the formation of Form B

Although any number of solvents may be used, including most lower alcohols, Form B is obtained in high yield preferably using methanol, which permits a high filtration rate of the crystalline material and allows the formation of Form B directly. The solubility of both the free base and L-tartaric acid are higher in methanol than in other lower alkyl alcohols.

The rate of formation of Form B may also be accelerated by employing the specific order of addition wherein the 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene free base is added to the solution of L-tartaric acid. To maximize the virtual concentration of L-tartaric acid present in the reaction, the methanolic solution of free base may be added to a solution containing either 1.1 or more equivalents of L-tartaric acid at 20° C. The desired anhydrous Form B may then be isolated directly and the polymorph conversion completed in less than 2 hours.

One optimized procedure for making the anhydrous Form B comprises charging a speck-free vessel with between 1.1 and 2.2 equivalents of L-tartaric acid and methanol (4 to 50 volumes), and stirring this mixture until dissolved and speck-free filtering the resulting solution into a crystallization vessel. 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]hexadeca-2(11),3,5,7,9-pentaene free base (1.0 equivalents) and methanol (4 to 50 volumes) are stirred in a vessel until dissolved at 0 to 50° C., more preferably at 20 to 25° C. The resulting solution of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene free base is then added over about a period of time ranging from 1 minute to 2 hours, more preferably over about 30 minutes, to the L-tartaric acid solution. The product was allowed to stir at 0 to 40° C., more preferably at 20 to 25° C., for between 1 and 48 hours, more preferably for about 1 hour, and then isolated by filtration. The product is dried generally under vacuum at 20 to 60° C., more preferably at 35 to 45° C., to give Form B of the L-tartrate salt of 5,8,14triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-

Both anhydrous Forms A and B can be converted to the monohydrate Form C by exposing either to a relative humidity (RH) of 100% or slurrying either of them in water. Form C is most readily obtained from either of Forms A or B by dissolving either in water at 20 to 50° C. followed by addition of an organic solvent in which the salt is not soluble, preferably methanol, ethanol or acetonitrile, and allowing the mixture to stir for between 1 and 30 minutes, preferably about 10 minutes. Upon filtering off the Form C which precipitates out as a white salt, the Form C salt may be air dried.

Noteworthy is that when exposed to conditions of 100% RH, Form B will convert to Form C within 2 days. Conversely, however, Form C readily converts to Form B upon exposure to 0% relative humidity conditions in roughly the same period of time. Hydrate Form C will however more

slowly dehydrate upon exposure to conditions of less than 50% RH. Experiments at 23% and 43% RH have verified this phenomena. Nonetheless, both Forms B and C appear to be relatively stable over a several month period at RH greater than 60%, as experiments at 75% and 87% relative 5 humidity have shown.

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Further, Form A can be obtained from Form C by dissolving Form C in a hot organic solvent, preferably ethanol, at or near its reflux point, preferably at about 75° C., and allowing it to stir for from 10 minutes to 3 hours, preferably 30 minutes. Hot filtering the mixture allows the collection of crystals which upon drying in a vacuum oven at 45° C. yields Form A.

The D-tartrate salt of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene has three polymorphs (Forms A, B' and C'), which exhibit the same x-ray diffraction characteristics, hygroscopicity, water content and thermal characteristics as the corresponding Forms A, B and C, respectively, of the L-tartrate salt; and are made in an identical manner as the corresponding L-tartrate salt polymorphs, with the exception that D-tartaric acid is employed in those procedures in place of L-tartraic acid.

The preparation of the anhydrous polymorph (Form X) of the D,L-tartrate salt of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene involves the steps of dissolving 5,8,14-triazatetracyclo ²⁵ [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent, preferably anhydrous ethanol, with about 1 to about 2.3 equivalents of D,L-tartaric acid, preferably 2.2 equivalents, at 20° C. to solvent reflux temperature for at least 2 hours, more preferably for at least 12 hours, most 30 preferably at least 24 hours; collecting the crystals formed, washing the product with solvent and air drying it. The hydrate polymorph (Form Y) of the D,L-tartrate salt may be made in an analogous fashion but with the use of a solvent admixed with water, preferably an ethanol and water 35 mixture, more preferably 20% aqueous ethanol. In addition, the meso-tartrate may be made in an analogous fashion to the D.L-tartrate.

Differential Scanning Calorimetry

The solid state thermal behavior of Forms A, B and C of the L-tartrate salt of 5,8,14-triazatetra-cyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene were investigated by differential scanning calorimetry (DSC). The traces for Forms A, B and C are shown in FIGS. 9A, 9B and 9C, respectively. The DSC thermograms were obtained on a Mettler Toledo DSC 821^e (STAR^e System). Generally, samples between 1 and 10 mg were prepared in crimped aluminum pans with a small pinhole. The measurements were run at a heating rate of 5° C. per minute in the range of 30 to 300° C.

As seen in FIG. 9A, the L-tartrate salt Form A exhibits an onset of melt transition at 223° C. with a melting peak accompanied by decomposition at 225° C. measured at a rate of 5° C. per minute. As seen in FIG. 9B, the L-tartrate salt Form B exhibited an onset of melt transition at 215° C. with a melting peak accompanied by decomposition at 218° C. measured at a rate of 5° C. per minute. As seen in FIG. 9C, the L-tartrate salt hydrate Form C exhibits a solid-solid transition onset at 73° C. with a peak at 76° C. This solid-solid transition is believed to correspond to the loss of water from the crystal lattice. A melt transition onset is also observed at 220° C., with a peak at 223° C. accompanied by decomposition.

The solid state thermal behavior of Forms X and Y of the D,L-tartrate salt of 5,8,14-triazatetra-cyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene were also investigated by DSC. As seen in FIG. 11A, the D,L-65 tartrate salt Form X (anhydrous) exhibits an onset of melting transition at 212° C. In FIG. 11B, the differential scanning

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calorimetric trace for the D,L-tartrate salt Form Y indicates an exhibits a solid-solid transition onset at 131° C. with a peak at 137° C. This solid-solid transition is believed to correspond to or to be associated with the loss of water from the crystal lattice. A melt transition onset for Form Y is also observed at 217° C. and is accompanied by decomposition.

One of skill in the art will however note that in DSC measurements there is a certain degree of variability in actual measured onset and peak temperatures which is dependant on rate of heating, crystal shape and purity, and a number of measurement parameters.

Powder X-ray Diffraction Patterns

The powder x-ray diffraction patterns for both Forms A, B and C of the L-tartrate salt were collected using a Bruker D5000 diffractometer (Bruker AXS, Madison, Wis.) equipped with copper radiation (CuK_{α}), fixed slits (1.0, 1.0, 0.6 mm), and a Kevex solid state detector. Data was collected from 3.0 to 40.0 degrees in two theta (20) using a step size of 0.04 degrees and a step time of 1.0 seconds.

The x-ray powder diffraction pattern of the L-tartrate salt Form Awas conducted with a copper anode with wavelength 1 at 1.54056 and wavelength 2 at 1.54439 (relative intensity: 0.500). The range for 2θ was between 3.0 to 40.0 degrees with a step size of 0.04 degrees, a step time of 1.00, a smoothing width of 0.300 and a threshold of 1.0.

The diffraction peaks at diffraction angles (20) in a measured powder X-ray diffraction analysis for the Form A are shown in Table I. The relative intensities, however, may change depending on the crystal size and morphology. The actual measured powder diffractogram is displayed in FIG.

TABLE I

Powder X-ray Diffraction Pattern for L-Tartrate Form A with Intensities and Peak Locations of Diffraction Lines.				
Angle 2θ	d-value (Å)	I (rel.)		
6.1	14.5	73.3		
11.8	7.5	6.1		
12.2	7.2	15.8		
13.0	6.8	23.9		
14.7	6.0	14.6		
16.8	5.3	99.5		
17.6	5.0	11.7		
18.3	4.8	7.0		
19.0	4.7	14.4		
19.4	4.6	28.4		
20.6	4.3	16.8		
21.9	4.1	100.0		
22.6	3.9	9.1		
23.9	3.7	13.4		
24.6	3.6	29.2		
27.2	3.3	10.5		
27.7	3.2	6.1		
28.8	3.1	8.0		
29.4	3.0	5.3		
29.8	3.0	15.9		
30.8	2.9	5.6		
32.0	2.8	5.8		
32.5	2.8	8.9		
34.0	2.6	6.0		
34.8	2.6	6.9		
35.2	2.5	8.8		
37.0	2.4	6.9		
37.5	2.4	8.6		
38.2	2.4	6.5		

Table II sets forth the 2θ , d-spacings and relative intensities representative of Form A. The numbers as listed are computer-generated.

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TABLE IV

TABLE II

	nd Peak Locations Report L-Tartrate Form A.	oresentative	5		nd Peak Locations Report L-Tartrate Form B.	presentative
Angle 2θ	d-value (Å)	I (rel.)		Angle 2θ	d-value (Å)	I (rel.)
6.1	14.5	73.3		5.9	15.0	57.0
12.2	7.2	15.8	10	12.8	6.9	27.2
				14.4	6.1	23.2
13.0	6.8	23.9		15.3	5.8	4.9
14.7	6.0	14.6		16.9	5.2	41.8
16.8	5.3	99.5		17.2	5.2	49.3
19.4	4.6	28.4		21.8	4.1	100.0
21.9	4.1	100.0	15	23.8	3.7	26.9
24.6	3.6	29.2		25.1	3.5	15.8

The x-ray powder diffraction pattern of the salt Form B ²⁰ was measured with the same equipment and under that same parameters used above for the measurement of Form A. The diffraction peaks at diffraction angles (2θ) in a measured powder X-ray diffraction analysis for the Form B are shown in Table III. Again, the relative intensities, however, may change depending on the crystal size and morphology. The actual measured powder diffractogram is displayed in FIG. 2.

The x-ray powder diffraction pattern of the salt Form C was measured with the same equipment and under that same parameters used above for the measurement of Form A. The diffraction peaks at diffraction angles (20) in a measured powder X-ray diffraction analysis for the Form C are shown in Table V. Again, the relative intensities, however, may change depending on the crystal size and morphology. The actual measured powder diffractogram is displayed in FIG. 3.

TABLE V

TABLE III

	ction Pattern for L-Ta Peak Locations of Dif				ction Pattern for L-Ta Peak Locations of Diff	
Angle 20	d-value (Å)	I (rel.)	35	Angle 20	d-value (Å)	I (rel.)
5.9	15.0	57.0		5.9	15.1	85.5
11.7	7.5	8.2		11.8	7.5	49.4
12.8	6.9	27.2		13.1	6.8	14.4
14.4	6.1	23.2	40	14.5	6.1	9.2
15.3	5.8	4.9		16.5	5.4	97.4
16.4	5.4	23.0		17.5	5.1	10.0
16.9	5.2	41.8		18.8	4.7	7.0
17.2	5.2	49.3		20.3	4.4	8.2
17.8	5.0	6.8		21.2	4.2	100.0
18.7	4.7	5.6	45	23.1	3.8	35.0
19.1	4.6	11.1	45	23.8	3.7	78.5
20.7	4.3	6.3		26.1	3.4	11.6
21.1	4.2	6.0		26.5	3.4	65.8
21.8	4.1	100.0		27.0	3.3	9.6
23.8	3.7	26.9		27.9	3.2	5.8
24.3	3.7	10.5		28.9	3.1	9.5
25.1	3.5	15.8	50	29.3	3.0	27.3
25.8	3.4	11.4		29.9	3.0	33.0
26.9	3.3	6.6		31.3	2.9	6.7
27.8	3.2	8.7		31.6	2.8	9.0
29.1	3.1	8.6		32.1	2.8	8.7
29.7	3.0	4.9		33.5	2.7	5.9
31.9	2.8	11.9	55	35.8	2.5	10.0
34.6	2.6	7.2		36.0	2.5	13.0
34.9	2.6	5.5		37.0	2.4	5.7
35.6	2.5	5.0		37.9	2.4	11.5
37.3	2.4	5.4		_	_	
38.8	2.3	5.4		_	_	_
_		_	60	_	_	
_	_				_	

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Table IV sets forth the 2θ , d-spacings, and relative intensities representative of Form B. The numbers as listed are computer-generated.

Table VI sets forth the 2θ , d-spacings, and relative intensities representative of Form C. The numbers as listed are computer-generated.

TABLE VI

	of L-Tartrate Form C.	
Angle 2θ	d-value (Å)	I (rel.)
5.9	15.1	85.5
11.8	7.5	49.4
16.5	5.4	97.4
21.2	4.2	100.0
23.1	3.8	35.0
23.8	3.7	78.5
26.5	3.4	65.8

As shown in FIG. 6, the overlay of the observed x-ray powder diffraction patterns for L-tartrate salt Forms A, B and C shows some x-ray powder diffraction peak shifting and that each Form has a distinctive powder pattern finger-print.

The x-ray powder diffraction pattern of the D,L-tartrate salt Form X (anhydrous) was measured with the same equipment and under that same parameters used above for the measurement of Form A, L-tartrate salt. The diffraction peaks at diffraction angles (2θ) in a measured powder X-ray diffraction analysis for the Form X are shown in Table VII. Again, the relative intensities, however, may change depending on the crystal size and morphology. The actual measured powder diffractogram is displayed in FIG. 10A.

TABLE VII

Powder X-ray Diffraction Pattern for D,L-Tartrate Form X

	Offraction Lines.	Peak Locations of D	with Intensities and
35	I (rel.)	d-value (Å)	Angle 20
_	100.0	14.6	6.0
	3.8	8.1	10.9
	13.0	7.7	11.5
	38.0	7.4	11.9
40	18.4	6.5	13.6
40	8.8	6.3	14.1
	27.6	5.9	15.0
	49.2	5.2	17.1
	10.3	4.8	18.3
	4.8	4.8	18.7
	6.0	4.5	19.6
45	49.5	4.0	22.1
	24.5	3.6	24.5
	4.3	3.5	25.3
	3.9	3.5	25.6
	11.8	3.4	26.4
	3.7	3.2	27.5
50	4.4	3.2	28.2
	11.7	2.8	31.8
	4.0	2.4	37.2
	3.7	2.4	37.3

Table VIII sets forth the 2θ , d-spacings, and relative 55 intensities representative of Form X. The numbers as listed are computer-generated.

TABLE VIII

	d Peak Locations Rep D,L-Tartrate Form X	
Angle 2θ	d-value (Å)	I (rel.)
6.0 11.9	14.6 7.4	100.0 38.0

TABLE VIII-continued

5	Intensities and Peak Locations Representative of D,L-Tartrate Form X.					
	Angle 20	d-value (Å)	I (rel.)			
10	15.0	5.9	27.6			
	17.1	5.2	49.2			
	22.1	4.0	49.5			
15	24.5	3.6	24.5			

The x-ray powder diffraction pattern of the D,L-tartrate salt Form Y (hydrate) was measured with the same equipment and under that same parameters used above for the measurement of Form A, L-tartrate salt. The diffraction peaks at diffraction angles (20) in a measured powder X-ray diffraction analysis for the Form Y are shown in Table IX.

25 Again, the relative intensities, however, may change depending on the crystal size and morphology. The actual measured powder diffractogram is displayed in FIG. 10B.

TABLE D

	Powder X-ray Diffraction Pattern for D,L-Tartrate Form Y with Intensities and Peak Locations of Diffraction Lines.						
35	Angle 20	d-value (Å)	I (rel.)				
	4.1	21.4	5.2				
	6.2	14.2	100.0				
	10.9	8.1	7.8				
	11.5	7.7	23.1				
	12.0	7.4	39.1				
40	12.5	7.1	4.6				
	13.5	6.5	16.6				
	14.4	6.1	14.7				
	15.0	5.9	16.4				
	15.2	5.8	32.7				
	15.6	5.7	9.6				
45	17.3	5.1	18.6				
73	18.1	4.9	32.2				
	18.7	4.7	7.1				
	19.9	4.5	24.7				
	21.1	4.2	7.0				
	21.7	4.1	11.0				
# 0	22.5	4.0	5.4				
50	23.2	3.8	12.2				
	24.0	3.7	52.7				
	25.1	3.5	75.1				
	25.5	3.5	10.3				
	26.1	3.4	8.5				
	27.5	3.2	17.9				
55	29.3	3.0	7.4				
	29.7	3.0	8.4				
	30.3	2.9	11.7				
	31.5	2.8	17.4				
	35.8	2.5	6.4				
	36.7	2.4	4.5				
60	37.3	2.4	4.6				
	39.1	2.3	5.4				

Table X sets forth the 2θ , d-spacings and relative intensities of Form Y. The numbers as listed are computergenerated.

		_	
TΛ	ы	- 14	V

	Intensities and Peak Locations Representative of D,L-Tartrate Form Y.				
Angle 20	d-value (Å)	I (rel.)			
6.2	14.2	100.0			
12.0	7.4	39.1			
15.2	5.8	32.7			
18.1	4.9	32.2			
24.0	3.7	52.7			
25.1	3.5	75.1			

Single Crystal X-ray Analysis

Single crystals for the L-tartrate salt Forms B and C were obtained and investigated by X-ray diffraction. For each form, a representative crystal was surveyed and a 1Å data set (maximum sin ⊕/λ=0.5) was collected on a Siemens R4RA/v diffractometer. Atomic scattering factors were taken from the *International Tables for X-Ray Crystallography*, Vol. IV, pp. 55, 99 and 149 (Birmingham: Kynoch Press, 1974). Single crystal X-ray data were collected at room temperature. All crystallographic calculations were facilitated by the SHELXTL™ system (SHELXTL™ Reference Manual, Version 5.1, Bruker AXS, Madison, Wis. 1997). The pertinent crystal data collection and refinement are summarized in Table XI below for Form B and in Table XII below for Form C.

For both Forms, the trial structure was obtained by direct methods and was then refined routinely. A difference map revealed two waters of crystallization—one for each salt molecule. Hydrogen positions were calculated wherever possible. The hydrogens on nitrogen and oxygen were located by difference Fourier techniques. The hydrogen parameters were added to the structure factor calculations but were not refined. The shifts calculated in the final cycles of least squares refinement were all less than 0.1 of the corresponding standard deviations. For Form B, the final R-index was 3.25%. For Form C, the final R-index was 3.47%. A final difference Fourier revealed no missing or misplaced electron density. The refined structure was plotted using the SHELXTL plotting package and is shown in FIGS. 8A (Form B) and 8B (Form C). The absolute configuration was based on the use of L(+)-tartaric acid.

Table XIII sets forth the atomic coordinates $(x10^4)$ and equivalent isotropic displacement parameters $(\mathring{A}^2 \times 10^3)$ for Form B. Table XIV lists the observed bond lengths $[\mathring{A}]$ and angles [°] for Form B. In Table XV, the anisotropic displacement parameters $(\mathring{A}^2 \times 10^3)$ for Form B are set forth to allow calculation of the anisotropic displacement factor exponent which has the form: $-2\pi^2[h^2 \ a^{*2}U_{11}+\ldots+2\ h\ k\ a^* \ b^*U_{12}]$. Finally, in Table XVI, below, hydrogen coordinates $(\times 10^4)$ and isotropic displacement parameters $(\mathring{A}^2 \times 10^3)$ for Form B are listed.

Table XVII sets forth the atomic coordinates ($\times 10^4$) and equivalent isotropic displacement parameters ($\mathring{A}^2 \times 10^3$) for Form C. Table XVIII lists the observed bond lengths [\mathring{A}] and 6 angles [°] for Form C. In Table XIX, the anisotropic displacement parameters ($\mathring{A}^2 \times 10^3$) for Form C are set forth to allow calculation of the anisotropic displacement factor exponent which has the form: $-2\pi^2[\mathring{h}^2 \ a^{*2}U_{11}+\ldots+2\ h\ k\ a^*$ b* U_{12}]. Finally, in Ta below, hydrogen Coordinates ($\times 10^4$) 6 and isotropic displacement parameters ($\mathring{A}^2 \times 10^3$) for Form C are listed.

TABLE XI

5	Crystal Structure Data and Measurement Parameters: L-Tartrate Salt Form B				
	Parameter		L-Tartrate Form B		
	Empirical formul	a	$C_{13}H_{14}N_3^+C_4H_5O_6^-$		
10	Formula weight		361.35		
	Crystal System		Orthorhombic		
	Space Group		P2(1)2(1)2(1)		
	Crystal Size, mm	3	$0.01 \times 0.08 \times 0.10$		
	a		7.0753(5) Å		
15	b		7.7846(5) Å		
	c		29.870(2) Å		
	α		90°		
	γ		90°		
	β		90°		
20	Volume		1645.21(19) Å ³		
	Density calc'd, p		1.459 g/cm ³		
	Z		4		
	Temperature		298(2) K		
25	Wavelength		1.54178 Å		
23	Absorption coeffi	cient	0.944 mm ⁻¹		
	F(000)		760		
	Reflections collec	eted	3490		
	Independent refle	ctions	1318 [R(int) = 0.0542]		
30	Refinement meth	od	Full-matrix least-squares on F2		
	Data/restraints/pa	rameters	1318/0/251		
	Goodness-of-fit of	n F ²	0.856		
	Final R indices [[> 2sigma(I)]	R1 = 0.0325, $wR2 = 0.0638$		
	Absolute structur	e parameter	0.0031(3)		
35	Largest diff. peak	and hole	0.115 and -0.150 e.Å-3		

TABLE XII

Crystal Structure Data and Measurement
Parameters: L-Tartrate Salt Form C

	Parameter	L-Tartrate Hydrate Form C
15	Empirical formula Formula weight	C ₁₃ H ₁₄ N ₃ ⁺ C ₄ H ₅ O ₆ ⁻ .H ₂ O 379.37
	Crystal System	Monoclinic
	Space Group	P2(1)
	Crystal Size, mm ³	$0.04 \times 0.38 \times 0.30$
	X-ray Code	F611
	a	7.5120 Å
50	b	29.854 Å
	c	7.671 Å
	α	90°
	γ	90°
	β	90.40°
	Volume	1720.3 Å ³
55	Density calc'd, ρ	1.465 g/cm ³
, ,	Z	4
	Temperature	298(2) K
	Wavelength	1.54178 Å
	Absorption coefficient	0.974 mm^{-1}
	F(000)	800
60	Reflections collected	1983
)U	Independent reflections	1817 [R(int) = 0.0224]
	Refinement method	Full-matrix least-squares on F ²
	Data/restraints/parameters	1817/0/528
	Goodness-of-fit on F ²	1.028
	Final R indices [I > 2sigma(I)]	R1 = 0.0347, $wR2 = 0.0834$
	Absolute structure parameter	0.0(3)
55	Largest diff. peak and hole	$0.168 \text{ and } -0.230 \text{ e.Å}^{-3}$

TABLE XIII TABLE XV

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114.7(4)

Atomic Coordinates (x104) And Equivalent Isotropic Displacement
Parameters ($\mathring{A}^2 \times 10^3$) For Form B. U(eq) is defined as one third
of the trace of the orthogonalized U., tensor.

Parameters ($\mathring{A}^2 \times 10^3$) For Form B. U(eq) is defined as one third of the trace of the orthogonalized U_{ij} tensor.		5		The Aniso	tropic disp	lacement	factor expo	onent takes the base U_{12} .				
	x	y	z	U(eq)	_		U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
N(1)	8211(8)	10638(7)	12233(1)	61(1)	_	N(1)	63(4)	70(4)	50(3)	12(2)	-2(3)	8(3)
C(2)	8968(8)	9093(11)	12235(2)	72(2)		C(2)	54(4)	114(6)	49(4)	20(4)	-3(3)	8(5)
C(3)	8093(11)	7629(9)	12047(2)	75(2)	10	C(3)	79(5)	78(5)	66(4)	14(4)	-6(4)	30(5)
N(4)	6431(8)	7715(6)	11853(1)	64(1)		N(4)	78(4)	54(4)	60(3)	8(3)	-9(3)	13(3)
C(5)	5624(9)	9313(8)	11834(2)	50(1)		C(5)	65(4)	45(4)	39(3)	5(3)	-3(3)	6(4)
C(6)	6502(8)	10752(9)	12025(2)	49(1)		C(6)	41(4)	69(5)	36(3)	8(3)	-9(3)	1(4)
C(7)	5676(8)	12396(7)	11985(1)	48(1)		C(7)	51(4)	56(5)	38(3)	3(3)	-2(3)	-5(4)
C(8)	4007(8)	12557(6)	11762(2)	41(1)		C(8)	45(4)	41(4)	38(3)	4(3)	1(3)	-3(4)
C(9)	3107(7)	11097(7)	11572(1)	42(1)	15	C(9)	46(4)	40(4)	40(3)	12(3)	9(3)	-4(4)
C(10)	3890(8)	9495(7)	11605(1)	49(1)		C(10)	54(4)	52(5)	41(3)	8(3)	-5(3)	-14(4)
C(11)	2865(7)	14122(6)	11634(1)	44(1)		C(11)	49(3)	43(3)	38(3)	-1(3)	1(3)	-1(3)
C(12)	891(6)	13347(6)	11573(1)	53(1)		C(12)	45(4)	63(4)	50(3)	6(3)	7(3)	3(3)
C(13)	1397(7)	11686(6)	11315(1)	46(1)		C(13)	42(3)	49(3)	48(3)	11(3)	-3(3)	-4(3)
C(14)	3510(6)	14823(6)	11182(1)	43(1)		C(14)	43(3)	39(3)	46(3)	-3(3)	2(2)	-1(3)
N(15)	3597(5)	13405(5)	10838(1)	39(1)	20	N(15)	35(3)	41(3)	40(2)	7(2)	3(2)	-2(2)
C(16)	1962(6)	12183(5)	10838(1)	46(1)	20	C(16)	42(3)	51(3)	44(3)	6(3)	-4(3)	-2(3)
C(20)	7858(9)	6393(6)	10523(1)	37(1)		C(20)	48(4)	30(4)	33(3)	9(3)	10(3)	-6(4)
O(21)	9522(5)	6116(4)	10603(1)	47(1)		O(21)	30(2)	41(2)	68(2)	3(2)	-5(2)	7(2)
O(22)	6680(4)	5324(4)	10349(1)	47(1)		O(22)	44(2)	22(2)	73(2)	-5(2)	-2(2)	2(2)
C(23)	7033(6)	8162(5)	10623(1)	32(1)		C(23)	26(3)	28(3)	42(3)	0(2)	7(2)	0(3)
O(24)	5062(4)	8318(4)	10542(1)	44(1)	25	O(24)	33(2)	33(2)	68(2)	-10(2)	4(2)	1(2)
C(25)	8063(6)	9486(5)	10339(1)	31(1)	23	C(25)	35(3)	25(3)	32(3)	-7(2)	-1(2)	4(3)
O(26)	7763(4)	9176(4)	9873(1)	35(1)		O(26)	35(2)	32(2)	38(2)	-5(1)	3(2)	-1(2)
C(27)	7520(6)	11321(6)	10465(2)	35(1)		C(27)	22(3)	40(4)	42(4)	-7(3)	-8(3)	1(3)
O(28)	7065(4)	11655(4)	10852(1)	43(1)		O(28)	53(2)	36(2)	41(2)	-7(2)	2(2)	2(2)
O(29)	7681(4)	12417(4)	10148(1)	47(1)		O(29)	74(2)	27(2)	41(2)	5(2)	7(2)	4(2)
` /	` '	` ′	` /	• /								

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Anisotropic Displacement Parameters ($\mathring{A}^2 \times 10^3$) For Form B.

TABLE XIV

Bond lengths [Å] and angles [°] for L-Tartrate Form B.						
Bond Lengths						
N(1)—C(2)	1.316(6)	C(11)—C(12)	1.532(6)			
N(1)—C(6)	1.362(6)	C(12)—C(13)	1.547(6)			
C(2)—C(3)	1.413(7)	C(13)—C(16)	1.531(5)			
C(3)—N(4)	1.314(7)	C(14)—N(15)	1.510(5)			
N(4)—C(5)	1.370(6)	N(15)—C(16)	1.498(5)			
C(5)—C(10)	1.411(6)	C(20)—O(21)	1.221(5)			
C(5)—C(6)	1.403(7)	C(20)—O(22)	1.288(5)			
C(6)—C(7)	1.412(6)	C(20)—C(23)	1.525(6)			
C(7)—C(8)	1.361(6)	C(23)—O(24)	1.420(5)			
C(8)—C(9)	1.421(6)	C(23)—C(25)	1.521(5)			
C(8)—C(11)	1.511(6)	C(25)—O(26)	1.428(5)			
C(9)—C(10)	1.368(6)	C(25)—C(27)	1.526(6)			
C(9)—C(13)	1.504(6)	C(27)—O(28)	1.227(5)			
C(11)—C(14)	1.526(5)	C(27)—O(29)	1.281(5)			
	Bond A	Angles				
C(2)—N(1)—C(6)	115.0(5)	C(14)—C(11)—C(12)	107.9(3)			
N(1)— $C(2)$ — $C(3)$	123.9(5)	C(11)— $C(12)$ — $C(13)$	100.2(3)			
N(4)— $C(3)$ — $C(2)$	121.8(5)	C(9)—C(13)—C(16)	110.0(4)			
C(3)—N(4)—C(5)	116.0(5)	C(9)—C(13)—C(12)	100.8(4)			
N(4)—C(5)—C(10)	118.3(6)	C(16)—C(13)—C(12)	108.2(4)			
N(4)— $C(5)$ — $C(6)$	121.5(6)	N(15)—C(14)—C(11)	110.6(4)			
C(10)— $C(5)$ — $C(6)$	120.2(6)	C(16)— $N(15)$ — $C(14)$	115.7(3)			
N(1)—C(6)—C(5)	121.8(6)	N(15)—C(16)—C(13)	111.2(3)			
N(1)—C(6)—C(7)	117.8(6)	O(21)— $C(20)$ — $O(22)$	126.1(5)			
C(5)— $C(6)$ — $C(7)$	120.3(5)	O(21)—C(20)—C(23)	119.4(5)			
C(8)—C(7)—C(6)	119.0(5)	O(22)— $C(20)$ — $C(23)$	114.5(5)			
C(7)—C(8)—C(9)	120.7(5)	O(24)—C(23)—C(25)	108.5(3)			
C(7)—C(8)—C(11)	131.5(5)	O(24)—C(23)—C(20)	114.8(4)			
C(9)—C(8)—C(11)	107.7(4)	C(25)—C(23)—C(20)	108.6(3)			
C(10)—C(9)—C(8)	121.2(5)	O(26)—C(25)—C(23)	111.0(3)			
C(10)—C(9)—C(13)	129.8(5)	O(26)—C(25)—C(27)	111.2(3)			
C(8)—C(9)—C(13)	108.7(5)	C(23)—C(25)—C(27)	112.0(4)			
C(9)—C(10)—C(5)	118.6(5)	O(28)— $C(27)$ — $O(29)$	125.4(4)			
C(8)—C(11)—C(14)	110.7(4)	O(28)—C(27)—C(25)	119.8(4)			

101.6(4) O(29)—C(27)—C(25)

C(8)—C(11)—C(12)

TABLE XVI

Hydrogen Coordinates	(×104) And Isotropic	Displacement Parameters
	$(\mathring{A}^2 \times 10^3)$ For Form	B.

	x	y	z	U(eq)
H(2A)	10149	8958	12367	80
H(3A)	8710	6576	12062	80
H(7A)	6264	13354	12108	80
H(10A)	3292	8546	11480	80
H(11A)	2887	15004	11868	80
H(12A)	76	14092	11398	80
H(12B)	295	13097	11858	80
H(13A)	372	10840	11321	80
H(14A)	2636	15704	11082	80
H(14B)	4748	15344	11213	80
H(15A)	3600(70)	14000(60)	10578(14)	80
H(15B)	4860(70)	12850(60)	10867(14)	80
H(16A)	2302	11156	10672	80
H(16B)	894	12713	10688	80
H(23A)	7270	8427	10939	80
H(24A)	4680(70)	7400(60)	10401(15)	80
H(25A)	9419	9355	10397	80
H(26A)	6710(70)	9120(70)	9841(17)	80
H(29A)	7180(60)	13930(80)	10298(14)	80

TABLE XVII

Atomic Coordinates (x10⁴) And Equivalent Isotropic Displacement Parameters (Å 2 x 10³) For Form C. U(eq) is defined as one third of the trace of the orthogonalized Uij tensor.

_		x	у	z	U(eq)
	N(1)	-159(7)	10186(3)	-1642(7)	45(1)
	C(2)	-239(10)	10333(3)	-58(10)	52(2)
	C(3)	1241(10)	10446(3)	959(9)	50(2)
	N(4)	2878(7)	10415(3)	368(6)	42(1)
	C(5)	3033(8)	10257(3)	-1310(8)	33(2)
	C(6)	1520(7)	10141(3)	-2302(8)	30(2)

TABLE XVII-continued

TABLE XVIII-continued

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	TABLE XVII-continued				TABLE XVIII-continued				
	Atomic Coordinates ($\times 10^4$) And Equivalent Isotropic Displacement Parameters ($\mathring{A}^2 \times 10^3$) For Form C. U(eq) is defined as one third of				Bond lengths [Å] and angles [°] for L-Tartrate Form C.			m C.	
		the orthogonalized			5	C(12)—N(13)	1.501(7)	C(62)—N(63)	1.511(7)
			_	T T/>		N(13)—C(14) C(14)—C(15)	1.504(6) 1.525(8)	N(63)—C(64) C(64)—C(65)	1.508(6) 1.537(8)
	X	У	Z	U(eq)		C(15)—C(16)	1.528(8)	C(65)—C(66)	1.533(8)
C(7)	1723(7)	9967	-4007(7)	32(2)		C(1X)—O(2X)	1.216(6)	C(1Y)— $O(3Y)$	1.259(7)
C(8)	3381(7)	9902(3)	-4622(7)	25(1)		C(1X)—O(3X)	1.295(6)	C(1Y)—O(2Y)	1.254(7)
C(9)	4905(7)	10018(3)	-3648(7)	25(1)	10	C(1X)—C(4X)	1.518(7)	C(1Y)— $C(4Y)$	1.543(8)
C(10)	4759(8)	10194(3)	-2016(8)	36(2)		C(4X)— $O(5X)$	1.417(6)	C(4Y)— $O(5Y)$	1.424(6)
C(11)	6537(7)	9881(3)	-4655(7)	31(2)		C(4X)—C(6X)	1.517(8)	C(4Y)—C(6Y)	1.526(8)
C(12)	7003(7) 5380(6)	9395(3)	-4191(7)	33(2) 27(1)		C(6X)—O(7X) C(6X)—C(8X)	1.419(7)	C(6Y)—O(7Y) C(6Y)—C(8Y)	1.413(7)
N(13) C(14)	4292(7)	9102(3) 9171(3)	-4292(6) -5922(7)	29(1)		C(8X)— $C(8X)C(8X)$ — $O(10X)$	1.541(7) 1.240(7)	C(8Y)— $C(8Y)$	1.521(8) 1.219(6)
C(15)	4011(7)	9668(3)	-6277(7)	28(1)	15	O(OTT) O(OTT)	1.267(7)	C(8Y)— $O(9Y)$	1.306(7)
C(16)	5826(8)	9887(3)	-6550(8)	41(2)	13	-()	Bond Angles		
C(1X)	1541(7)	7444(3)	-5634(8)	23(1)				<u> </u>	
O(2X)	1182(4)	7444(2)	-7182(5)	36(1)		C(2)— $N(1)$ — $C(6)$	115.5(6)	C(52)—N(51)—	115.6(5)
O(3X)	361(5)	7474(2)	-4418(5)	38(1)		N(4) O(2) O(2)	124 4(7)	C(56)	102.4/6
C(4X)	3457(6) 3649(5)	7425(3) 7280(2)	-4997(7) -3247(5)	24(1) 32(1)		N(1)— $C(2)$ — $C(3)$	124.4(7)	N(51)—C(52)— C(53)	123.4(6)
O(5X) C(6X)	4282(7)	7881(3)	-5336(7)	25(1)	20	N(4)—C(3)—C(2)	122.2(6)	N(54)—C(53)—	123.6(6)
O(7X)	3348(4)	8230(2)	-4482(5)	28(1)		11(1) ((3) ((2)	122.2(0)	C(52)	123.0(0)
C(8X)	6296(7)	7900(3)	-4948(7)	22(1)		C(3)—N(4)—C(5)	115.6(5)	C(53)—N(54)—	116.0(5)
O(9X)	7172(5)	7560(2)	-5428(5)	37(1)		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	• •	C(55)	, ,
O(10X)	6935(5)	8241(2)	-4266(5)	35(1)		N(4)— $C(5)$ — $C(6)$	121.1(6)	N(54)—C(55)—	119.6(5)
O(1W)	3226(6)	7996(2)	-924(5)	37(1)	25	NT(1) O(5) O(10)	110.0(5)	C(60)	100 1/5
N(51)	3493(6)	6295(3)	3311(7)	43(1)	23	N(4)—C(5)—C(10)	119.0(5)	N(54)—C(55)—	120.4(5)
C(52) C(53)	3598(9) 2144(9)	6141(3) 6031(3)	4922(9) 5890(8)	47(2) 45(2)		C(6)—C(5)—C(10)	119.8(6)	C(56) C(60)—C(55)—	120.0(5)
N(54)	494(7)	6065(3)	5313(7)	43(1)		C(0)—C(3)—C(10)	119.8(0)	C(56)	120.0(3)
C(55)	289(8)	6228(3)	3651(7)	30(1)		N(1)—C(6)—C(5)	121.3(6)	N(51)—C(56)—	121.0(6)
C(56)	1799(7)	6340(3)	2642(8)	30(2)		- (-) -(-) -(-)	(-)	C(55)	
C(57)	1574(8)	6528(2)	950(8)	32(2)	30	N(1)—C(6)—C(7)	118.9(5)	N(51)—C(56)—	118.8(5)
C(58)	-95(8)	6593(3)	320(7)	27(1)				C(57)	
C(59)	-1609(7)	6472(2)	1339(7)	25(1)		C(5)— $C(6)$ — $C(7)$	119.9(5)	C(55)—C(56)—	120.1(5)
C(60)	-1436(7)	6295(3)	2965(9)	35(2)		C(0) C(7) C(0)	110.0/5	C(57)	110.0(5)
C(61) C(62)	-3249(8) -3717(7)	6621(3) 7097(3)	334(8) 850(7)	32(2) 33(2)		C(8)— $C(7)$ — $C(6)$	118.8(5)	C(58)—C(57)— C(56)	119.0(5)
N(63)	-2088(6)	7392(3)	720(6)	26(1)	25	C(7)—C(8)—C(9)	121.9(5)	C(57)—C(58)—	120.4(5)
C(64)	-1014(7)	7329(3)	-916(6)	29(1)	33	0(1) 0(0) 0(1)	121.5(0)	C(59)	120.1(0)
C(65)	-765(7)	6828(3)	-1308(7)	30(1)		C(7)—C(8)—C(15)	130.5(5)	C(57)—C(58)—	131.4(5)
C(66)	-2599(8)	6612(3)	-1564(7)	36(2)				C(65)	
C(1 Y)	-2999(7)	8598(3)	27(7)	26(1)		C(9)— $C(8)$ — $C(15)$	107.4(5)	C(59)—C(58)—	107.9(5)
O(2Y)	-3633(5)	8257(2)	745(5)	35(1)		0(10) 0(0) 0(0)	100.0(5)	C(65)	101.0(5)
O(3Y)	-3884(5)	8934(2)	-462(5)	34(1)	40	C(10)— $C(9)$ — $C(8)$	120.9(5)	C(60)—C(59)— C(58)	121.9(5)
C(4Y)	-986(6)	8611(3)	-356(7)	20(1)		C(10)—C(9)—C(11)	130.2(5)	C(60)—C(59)—	130.8(5)
O(5Y) C(6Y)	-53(4) -163(7)	8261(2) 9070(3)	523(5) -16(7)	28(1) 23(1)		0(10) 0(5) 0(11)	100.2(0)	C(61)	100.0(0)
O(7 Y)	-165(7) -328(5)	9219(2)	1725(5)	33(1)		C(8)—C(9)—C(11)	108.7(5)	C(58)—C(59)—	107.1(5)
C(8 Y)	-326(3) 1746(7)	9048(3)	-658(8)	24(1)				C(61)	
O(9 Y)	2954(5)	9023(2)	572(5)	36(1)	45	C(9)— $C(10)$ — $C(5)$	118.7(5)	C(59)—C(60)—	118.7(5)
O(10Y)	2085(5)	9039(2)	-2209(5)	37(1)	45	0(0) 0(11) 0(12)	100.0(5)	C(55)	100.0(5)
O(2W)	54(6)	8500(2)	4066(5)	39(1)		C(9)— $C(11)$ — $C(12)$	108.9(5)	C(59)—C(61)—	109.2(5)
						C(9)—C(11)—C(16)	101.6(5)	C(62) C(59)—C(61)—	102.4(5)
						C(3) C(11) C(10)	101.0(3)	C(66)	102.1(0)
						C(12)—C(11)—C(16)	107.9(5)	C(62)—C(61)—	109.8(5)
	-	TABLE XVIII			50	. , . , . ,	` ′	C(66)	• /
					•	N(13)—C(12)—C(11)	110.8(5)	N(63)—C(62)—	109.8(5)
Bond	lengths [Å] a	nd angles [°] for L	-Tartrate Form	C.		-/		C(61)	
		1.7 (1 /17)	2)		•	C(14)— $N(13)$ — $C(12)$	113.6(4)	C(64)—N(63)—	114.9(4)
	Bor	nd Lengths (Form (<u>~)</u>			N(13)—C(14)—C(15)	110.8(4)	C(62) N(63)—C(64)—	110.6(4)
N(1)—C(2)		1.294(8) N(51)-	-C(52)	1.320(8)		14(13)—C(14)—C(13)	110.6(4)	C(65)	110.0(4)
N(1)—C(6)			–C(56)	1.375(7)	55	C(16)—C(15)—C(14)	108.6(5)	C(58)—C(65)—	101.8(4)
C(2)—C(3)			–C(53)	1.365(9)			(-)	C(66)	
C(3)—N(4)		1.316(8) C(53)-	-N(54)	1.317(8)		C(16)—C(15)—C(8)	101.6(4)	C(58)—C(65)—	109.1(4)
N(4)— $C(5)$			-C(55)	1.373(8)				C(64)	
C(5)—C(6)		1 1 1	-C(60)	1.410(8)		C(14)— $C(15)$ — $C(8)$	109.8(4)	C(66)—C(65)—	108.9(5)
C(5)—C(10)		. , , ,	-C(56)	1.417(8)	60	C(15) C(16) C(11)	00.7(4)	C(64)	00.2(4)
C(6)—C(7) C(7)—C(8)			−C(57) −C(58)	1.424(8) 1.355(8)		C(15)—C(16)—C(11)	99.7(4)	C(65)—C(66)— C(61)	99.3(4)
C(8)—C(9)			–C(56) –C(59)	1.431(8)		O(2X)—C(1X)—O(3X)	123.7(5)	O(3Y)— $C(1Y)$ —	125.2(5)
C(8)—C(15)			-C(65)	1.514(8)		, , , , , , , , , , , , , , , , , , , ,	(3)	O(2Y)	(-)
C(9)—C(10)		1.362(8) C(59)-	-C(60)	1.360(8)		O(2X)—C(1X)—C(4X)	121.2(5)	O(3Y)—C(1Y)—	116.1(5)
C(9)—C(11)			-C(61)	1.515(8)	65	overn overn -v		C(4Y)	
C(11)—C(12)			-C(62)	1.518(9)	0.5	O(3X)—C(1X)—C(4X)	115.1(5)	O(2Y)— $C(1Y)$ —	118.7(5)
C(11)—C(16)		1.545(8) C(61)-	—C(66)	1.539(8)				C(4 Y)	

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TABLE XVIII-continued

Bond lengths [Å] and angles [°] for L-Tartrate Form C.						
O(5X)—C(4X)—C(6X)	113.4(4)	O(5Y)—C(4Y)—	112.3(4)			
O(5X)—C(4X)—C(1X)	114.0(4)	C(6Y) O(5Y)—C(4Y)—	111.8(4)			
C(6X)—C(4X)—C(1X)	107.5(4)	C(1Y) C(6Y)—C(4Y)—	112.7(4)			
O(7X)—C(6X)—C(4X)	112.0(4)	C(1Y) O(7Y)—C(6Y)—	114.1(4)			
O(7X)—C(6X)—C(8X)	111.8(4)	C(8Y) O(7Y)—C(6Y)—	113.9(4)			
C(4X)—C(6X)—C(8X)	113.7(4)	C(4Y) C(8Y)—C(6Y)—	106.7(4)			
O(10X)—C(8X)—O(9X)	125.6(5)	C(4Y) O(10Y)—C(8Y)—	123.7(5)			
O(10X)—C(8X)—C(6X)	119.3(5)	O(9Y) O(10Y)—C(8Y)—	121.4(5)			
O(9X)—C(8X)—C(6X)	115.1(5)	C(6Y) O(9Y)—C(8Y)—	114.9(5)			
		C(6 Y)				

TABLE XIX

Anisotropic Displacement Parameters (Å² × 10³) For Form C. (The Anisotropic displacement factor exponent takes the form: $-2\pi 2$ [h² a*²U₁₁ + . . . + 2 h k a* b* U₁₂]).

	U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
N(1)	42(4)	46(4)	46(4)	-8(3)	4(3)	0(3)
C(2)	53(5)	51(5)	52(5)	-5(4)	9(4)	3(4)
C(3)	63(5)	40(4)	49(4)	-2(4)	19(4)	11(4)
N(4)	59(4)	30(3)	37(3)	-8(3)	-7(3)	11(3)
C(5)	44(4)	19(3)	35(4)	1(3)	-8(3)	9(3)
C(6)	27(3)	25(4)	39(4)	1(3)	3(3)	3(3)
C(7)	30(4)	36(4)	30(4)	-1(3)	-10(3)	4(3)
C(8)	28(4)	27(3)	19(3)	1(2)	-4(3)	3(3)
C(9)	27(3)	20(3)	29(4)	4(3)	-9(3)	0(3)
C(10)	33(4)	32(4)	44(4)	-8(3)	-14(3)	-4(3)
C(11)	30(3)	26(4)	38(4)	0(3)	-1(3)	-6(3)
C(12)	22(3)	44(4)	34(3)	0(3)	0(3)	0(3)
N(13)	27(3)	32(3)	21(3)	1(2)	0(2)	1(2)
C(14)	26(3)	34(4)	27(3)	-4(3)	-11(3)	-1(3)
C(15)	24(3)	29(4)	30(3)	7(3)	-5(3)	-2(3)
C(16)	42(4)	41(4)	39(4)	5(3)	7(3)	-2(3)
C(1X)	23(3)	19(3)	28(4)	-1(3)	8(3)	1(3)
O(2X)	28(2)	56(3)	25(2)	-7(2)	-2(2)	-1(2)
O(3X)	19(2)	69(3)	26(2)	8(2)	5(2)	2(2)
C(4X)	19(3)	30(3)	24(3)	5(3)	-1(2)	1(3)
O(5X)	29(2)	34(2)	33(2)	5(2)	-5(2)	8(2)
C(6X)	20(3)	28(3)	26(3)	-1(3)	2(2)	1(3)
O(7X)	21(2)	25(2)	36(2)	-3(2)	5(2)	4(2)
C(8X)	21(3)	30(4)	16(3)	-2(3)	1(2)	5(3)
O(9X)	19(2)	43(3)	49(3)	-10(2)	-1(2)	4(2)
O(10X)	26(2)	35(3)	45(2)	-10(2)	-7(2)	-1(2)
O(1W)	28(2)	47(3)	35(2)	-9(2)	1(2)	-1(2)
N(51)	29(3)	47(4)	54(4)	7(3)	-3(3)	8(3)
C(52)	44(4)	46(4)	51(5)	11(4)	-9(4)	4(3)
C(53)	50(5)	48(4)	35(4) 37(3)	2(3) 4(3)	-4(3)	10(4) 8(3)
N(54) C(55)	53(4) 34(4)	40(3) 28(3)	27(3)	5(3)	5(3) 4(3)	3(3)
C(56)	28(4)	25(3)	36(4)	-5(3)	2(3)	2(3)
C(57)	30(4)	34(4)	32(4)	4(3)	7(3)	3(3)
C(58)	32(4)	24(4)	24(3)	-1(3)	5(3)	-1(3)
C(59)	22(3)	21(3)	33(4)	0(3)	1(3)	-2(3)
C(60)	25(3)	32(4)	49(4)	3(3)	10(3)	-3(3)
C(61)	26(3)	30(4)	40(4)	2(3)	-6(3)	-6(3)
C(62)	25(3)	35(4)	38(4)	4(3)	0(3)	-2(3)
N(63)	25(3)	27(3)	27(3)	-2(2)	5(2)	1(2)
C(64)	36(3)	33(4)	18(3)	2(3)	8(3)	1(3)
C(65)	35(3)	33(4)	21(3)	-5(3)	3(3)	6(3)
C(66)	42(4)	32(4)	33(4)	-6(3)	-6(3)	2(3)
C(1Y)	23(3)	38(4)	17(3)	-1(3)	-6(2)	0(3)
O(2Y)	21(2)	42(3)	43(2)	11(2)	5(2)	-2(2)
O(3Y)	19(2)	41(3)	44(3)	11(2)	3(2)	8(2)
C(4 Y)	18(3)	22(3)	21(3)	3(2)	-1(2)	4(3)

TABLE XIX-continued

Anisotropic Displacement Parameters ($\mathring{A}^2 \times 10^3$) For Form C.
(The Anisotropic displacement factor exponent takes the form:
$-2\pi 2 \int_{0}^{2} h^{2} a^{*2} U_{11} + + 2 h k a^{*} b^{*} U_{12} D_{12}$

		U_{11}	U_{22}	U_{33}	U_{23}	U_{13}	U_{12}
10	O(5Y)	21(2)	31(2)	30(2)	3(2)	-2(2)	4(2)
	C(6Y)	23(3)	30(3)	17(3)	4(3)	1(2)	7(3)
	O(7Y)	32(2)	37(3)	31(3)	-3(2)	6(2)	7(2)
	C(8Y)	23(3)	16(3)	33(4)	3(3)	-2(3)	-4(2)
	O(9Y)	19(2)	61(3)	27(2)	-9(2)	-6(2)	5(2)
	O(10Y)	28(2)	57(3)	24(2)	4(2)	6(2)	1(2)
	O(2W)	32(2)	50(3)	35(3)	7(2)	-2(2)	3(2)

TABLE XX

Hydrogen Coordinates ($\times 10^4$) And Isotropic Displacement Parameters ($\mathring{A}^2 \times 10^3$) For Form C.

20		\11 \ \ \ \ \ \ \	, , , , , , , , , , , , , , , , , , , ,		
20		x	у	z	U(eq)
	H(2)	-1359	10366	435	80
	H(3)	1066	10546	2094	80
	H(7)	732	9899	-4690	80
2.5	H(10)	5770	10272	-1377	80
25	H(11)	7541	10086	-4476	80
	H(12A)	7896	9284	-4990	80
	H(12B)	7499	9383	-3021	80
	H(13X)	5710(100)	8750(30)	-4290(90)	80
	H(13Y)	4660(100)	9130(30)	-3380(100)	80
	H(14A)	3147	9025	-5797 ´	80
30	H(14B)	4897	9035	-6903	80
	H(15)	3202	9720	-7264	80
	H(16A)	5715	10190	-6996	80
	H(16B)	6570	9712	-7324	80
	H(3XX)	-980(110)	7490(30)	-4900(90)	80
	H(4X)	4082	7208	-5730	80
35	H(5XX)	3350(100)	7550(30)	-2600(100)	80
	H(6X)	4144	7936	-6589	80
	H(7XX)	3230(100)	8210(30)	-3240(100)	80
	H(1WX)	2060(110)	8070(30)	-390(90)	80
	H(1WY)	4280(110)	8050(30)	-270(100)	80
	H(52)	4720	6106	5423	80
40	H(53)	2329	5927	7019	80
	H(57)	2559	6605	286	80
	H(60)	-2435	6220	3610	80
	H(61)	-4250	6416	511	80
	H(62A)	-4647	7211	87	80
	H(62B)	-4158	7101	2035	80
45	H(63X)	-2480(100)	7730(30)	650(90)	80
43	H(63Y)	-1300(100)	7360(30)	1730(100)	80
	H(64A)	141	7470	-772	80
	H(64B)	-1620	7471	-1889	80
	H(65)	16	6777	-2307	80
	H(66A)	-2509	6308	-2010	80
	H(66B)	-3358	6788	-2329	80
50	H(4Y)	-860	8553	-1607	80
	H(5YX)	-140(100)	8240(30)	1670(100)	80
	H(6Y)	-797	9286	-757	80
	H(7YX)	-100(110)	9020(30)	2280(100)	80
	H(9YX)	4230(110)	8990(30)	40(90)	80
	H(2WX)	1040(110)	8370(30)	4630(100)	80
55	H(2WY)	-990(110)	8380(30)	4830(100)	80

The powder X-ray diffraction patterns for Forms B and C were calculated from the respective single crystal data gathered for each L-tartrate salt form via the use of the SFOG and XPOW computer programs provided as part of the SHELXTL™ computer library. The calculated powder pattern for Form B is shown in FIG. 4A. The calculated powder pattern for Form C is shown in FIG. 4B.

A comparison of the observed Form B powder pattern and 65 the calculated pattern results are displayed in the overlaid powder X-ray diffraction pattern of FIG. **5A**. The lower pattern trace corresponds to the calculated powder pattern (from single crystal results) and the upper pattern corresponds to a representative experimental powder pattern. The general match between the two patterns indicates the agreement between powder sample and the corresponding single crystal structure.

A comparison of the observed Form C powder pattern and the calculated pattern results are displayed in the overlaid powder X-ray diffraction pattern of FIG. 5B. The lower pattern trace corresponds to the calculated powder pattern (from single crystal results) and the upper pattern corresponds to a representative experimental powder pattern. The general match between the two patterns indicates the agreement between powder sample and the corresponding single 15 crystal structure.

Solid State NMR

Forms A, B and C of the L-tartrate salt of 5,8,14- 20 triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene were characterized by solid state NMR techniques. Approximately 300 mg of a sample was tightly packed into 7 mm ZrO spinner. The ¹³C spectra were collected using cross-polarization magic angle spinning (CPMAS) at 295 K on Bruker 7 mm WB MAS probe positioned into a widebore Bruker Avance DRX 500 MHz NMR spectrometer. The samples were spun at 7 kHz. The cross-polarization contact time was set to 1 ms. The total of 512 scans were acquired for most of the samples resulting in approximately 30 minute acquisition times. The spectra were referenced using external sample of adamantane with the most upfield methyl signal set to 29.5 ppm.

The resulting ¹³C CPMAS spectra of Forms A, B and C are shown in FIGS. 7A, 7B and 7C, respectively. The samples behaved reasonably well from the point of view of solid state spectra quality. The resolution was good and the sensitivity was acceptable. The spectra features of all the compounds differ substantially from each other suggesting that solid state NMR can easily resolve the minor physical/chemical differences between the samples.

All the peaks marked with asterisks (*) are spinning sidebands in FIG. 7A, 7B and 7C. The spinning sidebands are displaced at multiple of the spinning frequencies along both sides of the real peaks (centerbands). The spinning speed was set to 7 kHz which at the 500 MHz magnet 50 translates into 55.7 ppm. The sideband intensities depend on the spinning speed (the higher the speed the lower the sideband intensity) and on the size of the anisotropic contribution of the chemical shielding for the given carbon. They can be easily distinguished from centerbands by variable spinning speed experiments. Carbonyl and aromatic sites tend to have very intense sidebands due to their large chemical shielding anisotropies. CH and CH₂ type of carbons give origin to relatively small spinning sidebands. Methyl groups (CH₃) usually don't generate any sidebands.

The major resonance peaks (those downfield from 100 ppm; ±0.1 ppm) for the solid state carbon spectrum of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 65 5,7,9-pentaene L-tartrate salt Forms A, B and C are listed in Table XXI.

TABLE XXI

Major Solid State ¹³C-NMR Resonance Peaks For 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene L-Tartrate Salt Forms A, B and C (Only Peaks Downfield from 100 ppm Listed) (Adamantane 29.5 ppm Standard).

	FORM A ¹³ C (ppm) Solid	FORM B ¹³ C (ppm) Solid	FORM C ¹³ C (ppm) Solid
, —	178.4	179.2	179.0
	149.3	178.0	176.1
	147.4	147.4	147.5
	145.1	145.2	144.5
	122.9	144.4	124.6
5		124.8	
•		122.5	

The L-tartrate, the D-tartrate, the D,L-tartrate and the meso-tartrate salts of the invention (hereafter "the active salts") can be administered via either the oral, transdermal (e.g., through the use of a patch), intranasal, sublingual, rectal, parenteral or topical routes. Transdermal and oral administration are preferred. These salts are, most desirably, administered in dosages ranging from about 0.01 mg up to about 1500 mg per day, preferably from about 0.1 to about 300 mg per day in single or divided doses, although variations will necessarily occur depending upon the weight and condition of the subject being treated and the particular route of administration chosen. However, a dosage level that is in the range of about 0.001 mg to about 10 mg per kg of body weight per day is most desirably employed. Variations may nevertheless occur depending upon the weight and condition of the persons being treated and their individual responses to said medicament, as well as on the type of pharmaceutical formulation chosen and the time period and interval during which such administration is carried out. In some instances, dosage levels below the lower limit of the aforesaid range may be more than adequate, while in other cases still larger doses may be employed without causing any harmful side effects, provided that such larger doses are first divided into several small doses for administration throughout the day.

The active salts can be administered alone or in combination with pharmaceutically acceptable carriers or diluents by any of the several routes previously indicated. More particularly, the active salts can be administered in a wide variety of different dosage forms, e.g., they may be combined with various pharmaceutically acceptable inert carriers in the form of tablets, capsules, transdermal patches, lozenges, troches, hard candies, powders, sprays, creams, salves, suppositories, jellies, gels, pastes, lotions, ointments, aqueous suspensions, injectable solutions, elixirs, syrups, and the like. Such carriers include solid diluents or fillers, sterile aqueous media and various non-toxic organic solvents. In addition, oral pharmaceutical compositions can be suitably sweetened and/or flavored. In general, the active compound is present in such dosage forms at concentration levels ranging from about 5.0% to about 70% by weight.

For oral administration, tablets containing various excipients such as microcrystalline cellulose, sodium citrate, calcium carbonate, dicalcium phosphate and glycine may be employed along with various disintegrants such as starch (preferably corn, potato or tapioca starch), alginic acid and certain complex silicates, together with granulation binders like polyvinylpyrrolidone, sucrose, gelatin and acacia. Additionally, lubricating agents such as magnesium stearate, sodium lauryl sulfate and talc can be used for tabletting purposes. Solid compositions of a similar type may also be

employed as fillers in gelatin capsules; preferred materials in this connection also include lactose or milk sugar, as well as high molecular weight polyethylene glycols. When aqueous suspensions and/or elixirs are desired for oral administration the active ingredient may be combined with various sweetening or flavoring agents, coloring matter and, if so desired, emulsifying and/or suspending agents, together with such diluents as water, ethanol, propylene glycol, glycerin and various combinations thereof.

For parenteral administration, a solution of an active salt 10 in either sesame or peanut oil or in aqueous propylene glycol can be employed. The aqueous solutions should be suitably buffered (preferably pH greater than 8), if necessary, and the liquid diluent first rendered isotonic. These aqueous solutions are suitable for intravenous injection purposes. The 15 oily solutions are suitable for intraarticular, intramuscular and subcutaneous injection purposes. The preparation of all these solutions under sterile conditions is readily accomplished by standard pharmaceutical techniques well known to those skilled in the art.

It is also possible to administer the active salts topically and this can be done by way of creams, a patch, jellies, gels, pastes, ointments and the like, in accordance with standard pharmaceutical practice.

EXAMPLES

The following examples illustrate the methods and compounds of the present invention. It will be understood, however, that the invention is not limited to the specific Examples.

Example 1

L-Tartrate Salt of 5,8,14-Triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (Anhydrous Polymorph, Form B)

A speck-free vessel was charged with L-tartaric acid (780 55 grams, 1.1 equiv.) and methanol (7.5 L). The contents of the vessel were stirred until solution and speck free filtered into the crystallization vessel. 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene free base (992 grams) and methanol (7.5 L) were dissolved in the 60 vessel; the mixture was maintained at between 20 to 25° C. The solution of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene free base was added over about 45 minutes to the L-tartaric acid solution through a filter to render the solution speck and fiber free. The product 65 was allowed to stir at 20 to 25° C. overnight and isolated by filtration. The product was dried under vacuum at 35 to 45°

C. to give 1618.4 grams (95.4%) of 5,8,14-triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene L-tartrate salt Form B (MW 361.36). M.p. 210.5° C.; verified as Form B by powder x-ray diffraction.

Example 2

L-Tartrate Salt of 5,8,14-Triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (Anhydrous Polymorph, Form A)

A reactor was charged with 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene free base (2 g; 0.0095 mole, 1.0 equiv.) and methanol (60 mL, 30 mL/g). The mixture was stirred at 20 to 25° C. until completely dissolved. A second reactor containing a solution of L-tartaric acid (1.55 g, 0.0103 mole, 1.1 equiv.) dissolved in methanol (60 mL, 30 mL/g) was heated to reflux in methanol (i.e., 60 to 66° C.). The free base solution was added to the L-tartaric acid solution at methanolic reflux temperature over 20 minutes. The resulting slurry was cooled to 20 to 25° C. over a 1 hour period. The reaction mixture was allowed to stir for approximately 2 hours followed by isolation of the product by filtration. The solid product was washed with methanol (10 mL), then dried under vacuum at 30 to 35° C. to give 3.3 grams (97%) of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3, 5,7,9-pentaene L-tartrate Form A. The identity as Form A was determined by PXRD as compared with standard samples.

Example 3

L-Tartrate Salt Form C of 5,8,14-Triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (Form C)

Preparation of CP-526,555-18 Form C from Form A or Form B:

L-tartrate salt Form B (~5 g) was dissolved in water (10 to 15 ml). Acetonitrile (200 to 300 ml) was added and Form C formed as a white precipitate. The resulting slurry was allowed to stir for 10 minutes and then filtered. The wet cake was then allowed to air dry. Product was determined to be Form C by NIR spectroscopy, DSC and PXRD analysis. This procedure may be run with Form A to yield Form C.

Example 4

L-Tartrate Salt Form A of 5,8,14-Triazatetracyclo [10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene (Form A)

Preparation of Form A from Form C: L-tartrate salt Form C (~2 g) was added to 200 to 300 mL hot ethanol (~75° C.) and allowed to stir for 30 minutes. The sample was filtered hot and then dried in a 45° C. vacuum oven (house vacuum). The material was determined to be Form A by NIR spectroscopy, DSC, and PXRD analysis.

What is claimed is:

- 1. The tartrate salt of 5,8,14-triazatetracyclo $[10.3.1.0^{2,11}.0^{4,9}]$ -hexadeca-2(11),3,5,7,9-pentaene.
- 2. A compound according to claim 1 which is the L-tartrate salt.
 - 3. A compound according to claim 2 which is anhydrous.
- 4. A compound according to claim 1 which is the D,L-tartrate salt.
 - 5. A compound according to claim 4 which is anhydrous.
- 6. A compound according to claim 1 which is D-tartrate

- 7. A compound according to claim 6 which is anhydrous.
- 8. A compound according to claim 6 which is a hydrate.
- A compound according to claim 1 which is the mesotartrate salt.
 - 10. The L-tartrate salt of claim 2 that is a hydrate.
- 11. A compound according to claim 10 where the hydrate is a monohydrate.
 - 12. A D,L-tartrate of claim 4 which is a hydrate.
- 13. A pharmaceutical composition comprising a pharmaceutically acceptable carrier and a compound according to 10 any of claims 1, 2, 6 or 9.
- 14. A method of treatment for nicotine dependency, addiction and withdrawal comprising the administration of a compound according to any of claims 1, 2, 6 or 9 to a subject in need thereof.
- 15. A process for the preparation of a compound according to claim 1 comprising the steps of
 - (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with between about 1 and about 2 equivalents of ²⁰ L-tartaric acid; and
 - (ii) collecting the crystals formed.
- 16. A process according to claim 15 wherein 1.1 equivalents of L-tartaric acid are employed and the tartaric acid is added to a solution containing the free base.
- 17. A process according to claim 15 wherein the contacting step is allowed to proceed above 45° C.
- **18**. A process according to claim **15** wherein the contacting step is allowed to proceed for less than 2 hours.
- 19. A process according to claim 15 wherein the suitable solvent is selected from the group consisting of an (C_1-C_6) alkyl alcohol, an (C_1-C_6) alkyl ketone, an (C_1-C_6) alkyl ether, acetonitrile and an (C_1-C_6) alkyl ester.
- **20.** A process according to claim **15** wherein the suitable solvent is ethanol or methanol.
- 21. A process for the preparation of a compound according to claim 1 comprising the steps of
 - (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with between about 1 and about 2.3 equivalents of L-tartaric acid; and
 - (ii) collecting the crystals formed.
- 22. A process according to claim 21 wherein 1.1 equivalents of L-tartaric acid are employed and the free base in 45 solution is added to a solution containing L-tartaric acid.
- 23. A process according to claim 21 wherein the contact step is allowed to proceed for at least 2 hours.
- **24.** A process according to claim **21** wherein the contact step is allowed to proceed for at least 12 hours.
- **25**. A process according to claim **21** wherein the suitable solvent is selected from the group consisting of an (C_1-C_6) alkyl alcohol, an (C_1-C_6) alkyl ketone, an (C_1-C_6) alkyl ether, acetonitrile and an (C_1-C_6) alkyl ester.

- 26. A process according to claim 21 wherein the suitable solvent is methanol or ethanol.
- 27. A process according to claim 21 wherein the suitable solvent is methanol.
- **28**. A process for the preparation of a compound according to claim **16** comprising the steps of
 - (i) contacting an anhydrous L-tartrate salt of 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5, 7,9-pentaene with water; and
 - (ii) collecting the crystals formed.
- 29. A process according to claim 28 wherein the contacting of step (i) comprises exposing the anhydrous L-tartrate salt to greater than 70% humidity.
- **30.** A process according to claim **28** wherein the contacting of step (i) comprises slurrying the anhydrous L-tartrate salt with water.
- 31. A process according to claim 28 wherein step (i) comprises the addition of an organic solvent.
- 32. A process according to claim 28 wherein step (i) comprises the addition of methanol, ethanol or acetonitrile.
- **33.** A process for the preparation of a compound according to claim **25** comprising the steps of
 - (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with about 1 to about 2.3 equivalents of D,L-tartaric acid; and
 - (ii) collecting the crystals formed.
- **34**. A process according to claim **33** wherein about 2.2 equivalents of D,L-tartaric acid is employed and the free base in solution is added to a solution containing D,L-tartaric acid.
- **35**. A process according to claim **33** wherein the contact step is allowed to proceed for at least 24 hours.
- **36**. A process according to claim **33** wherein the suitable solvent is anhydrous ethanol.
- 37. A process for the preparation of a compound according to claim 12 comprising the steps of
 - (i) contacting 5,8,14-triazatetracyclo[10.3.1.0^{2,11}.0^{4,9}]-hexadeca-2(11),3,5,7,9-pentaene in a suitable solvent with about 1 to about 2.3 equivalents of D,L-tartaric acid; and
 - (ii) collecting the crystals formed.
- **38**. A process according to claim **37** wherein about 2.2 equivalents of D,L-tartaric acid is employed and the free base in solution is added to a solution containing D,L-tartaric acid.
- 39. A process according to claim 37 wherein the contact step is allowed to proceed for at least 24 hours.
 - **40**. A process according to claim **37** wherein the suitable solvent is 20% aqueous ethanol.

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